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CLAIMS

(57)[Claim(s)]

(A) An organic polymer which has a hydroxyl group or a hydrofytic basis combined with a silicon atom, and has at least one silicon content group which can construct a bridge by forming a siloxane bond,

potassium, carboxylic acid barium, carboxylic acid manganese, a hardenability constituent containing B) Carboxylic acid calcium, carboxylic acid vanadium, carboxylic acid titanium, carboxylic acid one or more sorts of carboxylic acid metal salt chosen from carboxylic acid nickel

(C) The hardenability constituent according to claim 1 which becomes considering an amine compound as an essential ingredient as an ingredient.

Claim 3]

A) A number average molecular weight is within the limits of 500-50,000, and an organic polymer of an ingredient is a general formula to an end and/or a side chain of a main chain (1).

Formula 1

numbers 1-20. It is the Tori ORGANO siloxy group shown by the aryl group of the carbon numbers 6the inside of a formula, R^1 , and R^2 — respectively — independent — the alkyl group of the carbon 20, the aralkyl group of the carbon numbers 7-20, or (R') SiO- (R' is the substitution or the

nydroxyl group or a hydrolytic basis independently, respectively. a is 0, 1, 2, or 3, b is 0, 1, or 2, and a unsubstituted hydrocarbon group of the carbon numbers 1-20 independently, respectively). X is a and b are not simultaneously set to 0, m is an integer of 0, or 1-19.

The hardenability constituent according to claim 1 or 2 coming out and having one or more hydrolytic silyl groups per molecule expressed.

The hardenability constituent according to claim 3, wherein X is an alkoxy group.

Claim 5]

(A) A hardenability constituent given in any 1 paragraph of Claims 1-4 whose organic polymers of an ngredient are a polyoxyalkylene series polymer and/or a saturated hydrocarbon system polymer. Claim 6

hydrocarbon system polymer has a repeating unit resulting from isobutylene 50% of the weight or The hardenability constituent according to claim 5 which is a polymer, wherein said saturated nore in a total amount

hardenability constituent given in any 1 paragraph of Claims 1–6 to which carboxylic acid potassium, carboxylic acid manganese, and carboxylic acid nickel use as the main ingredients carboxylic acid metal salt expressed with general formula (2), (3), and (6) - (10), (B) Carboxylic acid calcium of an ingredient, carboxylic acid vanadium, carboxylic acid titanium, A

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respectively.

JP,4150220,B [CLAIMS]

Ca(OCOR) 2 (2)

Ti(0COR) 4 (6) V(OCOR) 3 (3)

K(0C0R) (7)

Ba(OCOR) 2 (8)

Mn(OCOR), (9)

nickel(OCOR) 2 (10)

(The inside R of a formula is substitution or an unsubstituted hydrocarbon group, and may include a carbon carbon double bond.)

salt of an ingredient is the carboxylic acid metal salt in which the melting point has an acid radical of (B) The hardenability constituent according to any one of claims 1 to 7 whose carboxylic acid metal carboxylic acid which is 65 ** or less.

(B) The hardenability constituent according to any one of claims 1 to 7 which is the carboxylic acid metal salt which has an acid radical of carboxylic acid whose carbon numbers in which carboxylic acid metal salt of an ingredient contains carbon of a carbonyl group are 2-17.

carboxylic acid group content compound in which carboxylic acid metal salt of an ingredient is chosen (B) A hardenability constituent given in any 1 paragraph of Claims 1-8 which are metal saft of a from octylic acid, 2–ethylhexanoic acid, neo decanoic acid, oleic acid, or naphthenic acid.

[Claim 11]

(A) A hardenability constituent given in any 1 paragraph of Claims 1–10 containing the (B) ingredient of quantity which serves as 0.005 – 5 weight section by metallic element conversion contained in the (B) ingredient to ingredient 100 weight section.

(A) A hardenability constituent given in any 1 paragraph of Claims 2–10 containing the (B) ingredient of quantity which serves as 0.005 – 5 weight section by metallic element conversion contained in the (B) ingredient, the (C) ingredient 0.01 – 20 weight sections to ingredient 100 weight section.

[Translation done.]

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DETAILED DESCRIPTION

Detailed Description of the Invention

Field of the Invention

In this invention, it has the hydroxyl group or hydrolytic basis combined with the silicon atom, and a siloxane bond is formed.

Therefore, it is related with the hardenability constituent containing the organic polymer which has a silicon content group (henceforth a "reactive silicon group") which can construct a bridge.

Description of the Prior Art]

It is known that the organic polymer which contains at least one reactive silicon group in a molecule accompanied by the hydrolysis reaction of a reactive silicon group, etc., and a rubber-like hardened has the interesting character in which construct a bridge by formation of the siloxane bond material is obtained with hygroscopic surface moisture etc. also in a room temperature.

polyisobutylene system polymer, It is indicated by JP,S52-73998,A, JP,H5-125272,A, JP,H3-72527,A, In the polymer which it has, these reactive silicon groups a polyoxyalkylene series polymer and a JP,S63-6003,A, JP,S63-6041,A, JP,H1-38407,A, JP,H8-231758,A, etc.

produced industrially, and are widely used for uses, such as a sealing material, adhesives, and a paint. Especially a polyoxyalkylene series polymer and a polyisobutylene system polymer are already

used for a structural sealing material, the sealing material for multiple glass, etc., it is effective. The neat resistance, a water resisting property, weatherability, etc., if the hardened material obtained is 50% of the weight or more in a total amount is preferred as a sealant aiming at moisture proof from especially the thing to excel in low moisture permeability and low gas permeability. isobutylene system polymer which has a repeating unit which furthermore originates in isobutylene When an organic polymer is a saturated hydrocarbon system polymer especially, since it excels in

flattery nature to the use part ranging from being used to a long period of time in order to give watertight and airtightness becomes very important, and revealing sufficient elongation and intensity Such a sealing material generally fills up the joined part and crevice between various members, the is called for.

bisacetylacetonate, is used widely, and when the movement of the part which places especially a sealing material is large, since the hardened material which has stability is obtained, many divalent tin purpose of a silanol condensation catalyst being used for bridge construction hardening of an organic condensation reaction. As such a silanol condensation catalyst, divalent tin, such as octylic acid (2-Amine compounds, such as lauryl amine, are used together as a co-catalyst in many cases for the ethylhexanoic acid) tin, The tin series catalyst represented by tetravalent tin, such as dibutyltin polymer which, on the other hand, has such a reactive silicon group, and also accelerating a

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...

JP,4150220,B [DETAILED DESCRIPTION]

elongation of a hardened material which are acquired may be insufficient, and an improvement of the compound of a co-catalyst and is used as a curing catalyst of the (A) ingredient, the intensity and However, when octylic acid tin which is divalent tin, for example is used together with the amine further physical properties is desired.

hardened material obtained has emollience, i.e., having the character to follow to the long-term stress catalyst of 1 liquid mold-curing nature constituent. However, since toxicity in case a small amount of Although most is a dibutyltin type, the latter tetravalent tin series catalyst being used now, While the change from the outside and fast curability, it is broadly used from the ability to apply as a curing tributyltin contains in this is regarded as questionable, development of the silanol condensation catalyst of a non-tin series is desired.

with carboxylate of tin. As an example in which the carboxylic aoid metal salt of non-tin was used for organopolysiloxane constituent, the carboxylic acid metal salt of various kinds of non-tin is indicated 12860,A) and carboxylic acid cerium (JP,2000-313814,A), there was no example broadly examined in the curing catalyst of the constituent which, on the other hand, contains the organic polymer which In JP,S35-2795,B, JP,S32-3742,B, JP,S35-9639,B, JP,S37-3271,B, etc., As a curing catalyst of an has a reactive silicon group, Although there were carboxylic acid bismuth (JP,H5-39428,A, JP,H9various carboxylic acid metal salt until now.

[Problem(s) to be Solved by the Invention]

The purpose of this invention is as follows.

Have practical hardenability and stability with a good hardened material obtained.

Provide a hardenability constituent useful as the structural sealing material asked especially for high durability, the sealing material for multiple glass, the moisture seal material used for an electric electronic component, adhesives, etc. from revealing the outstanding intensity and elongation.

Means for Solving the Problem

oarboxylic acid cobalt, and a carboxylic acid zirconium, Securing moderate hardenability and stability, oatalyst of the (A) ingredient Carboxylic acid calcium, Carboxylic acid vanadium, carboxylic acid iron. In order that this invention persons may solve such a problem, as a result of inquiring, as a curing it finds out revealing high intensity and the physical properties of high elongation compared with a carboxylic acid titanium, carboxylic acid potassium, By using one or more sorts of carboxylic acid metal salt chosen from carboxylic acid barium, carboxylic acid manganese, carboxylic acid nickel, case where octylic acid tin is used as a curing catalyst, and came to complete this invention

forming a siloxane bond, (B) It is related with a hardenability constituent containing one or more sorts Namely, this invention has a hydroxyl group or a hydrolytic basis combined with the (A) silicon atom. An organic polymer which has at least one silicon content group which can construct a bridge by of carboxylic acid metal salt chosen from carboxylic acid calcium, carboxylic acid vanadium, carboxylic acid titanium, carboxylic acid potassium, carboxylic acid barium, carboxylic acid manganese, and carboxylic acid nickel.

An amine compound is related with the aforementioned hardenability constituent which becomes as an essential ingredient as a (C) ingredient. As a desirable embodiment, a number average molecular weight is within the limits of 500-50,000, and an organic polymer of the (A) ingredient is a general formula to an end and/or a side chain of a main

[Formula 2]

.

numbers 1-20. It is the Tori ORGANO siloxy group shown by the aryl group of the carbon numbers 6hydroxyl group or a hydrolytic basis independently, respectively. a is 0, 1, 2, or 3, b is 0, 1, or 2, and a and b are not simultaneously set to 0 m -- the integer of 0, or 1-19 -- it is -- it is related with a hardenability constituent given in said either having one or more hydrolytic silyl groups per molecule the inside of a formula, R¹, and R² — respectively — independent — the alkyl group of the carbon unsubstituted hydrocarbon group of the carbon numbers 1-20 independently, respectively). X is a 20, the aralkyl group of the carbon numbers 7–20, or (R) $_3{\rm SiO}-({\rm R}'$ is the substitution or the expressed.

It is related with the aforementioned hardenability constituent characterized by X being an alkoxy group as a desirable embodiment.

As a desirable embodiment, an organic polymer of the (A) ingredient is related with a hardenability constituent given in said either which is a polyoxyalkylene series polymer and/or a saturated hydrocarbon system polymer.

aforementioned hardenability constituent which is a polymer having a repeating unit resulting from As a desirable embodiment, said saturated hydrocarbon system polymer is related with the isobutylene 50% of the weight or more in a total amount.

manganese, and carboxylic acid nickel are related with a hardenability constituent given in said either which uses as the main ingredients carboxylic acid metal salt expressed with general formula (2), (3), As a desirable embodiment, carboxylic acid calcium of the (B) ingredient, Carboxylic acid vanadium, carboxylic acid titanium, carboxylic acid potassium, Carboxylic acid barium, carboxylic acid and (6) – (10), respectively.

Ca(OCOR) 2 (2)

V(OCOR) 3 (3)

[(OCOR) 4 (6) K(OCOR)(7)

Ba(OCOR) 2 (8)

Mn(OCOR) 2 (9)

 $nickel(OCOR)_2$ (10)

The inside R of a formula is substitution or an unsubstituted hydrocarbon group, and may include a carbon carbon double bond.)

hardenability constituent given in said either which is the carboxylic acid metal salt in which the melting point has an acid radical of carboxylic acid which is 65 ** or less. As a desirable embodiment, carboxylic acid metal salt of the (B) ingredient is related with a

As a desirable embodiment, a carbon number in which carboxylic acid metal salt of the (B) ingredient oontains carbon of a carbonyl group is related with a hardenability constituent given in said either which is the carboxylic acid metal salt which has an acid radical of carboxylic acid which is 2-17.

nardenability constituent given in said either which is metal salt of a carboxylic acid group content As a desirable embodiment, carboxylic acid metal salt of the (B) ingredient is related with a

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JP,4150220,B [DETAILED DESCRIPTION]

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compound chosen from octylic acid, 2-ethylhexanoic acid, neo decanoic acid, oleic acid,

naphthenic acid.

quantity which serves as 0.005 - 5 weight section by metallic element conversion contained in the It is related with a hardenability constituent given in said either containing the (B) ingredient of (B) ingredient to (A) ingredient 100 weight section as a desirable embodiment.

quantity which serves as 0.005 - 5 weight section by metallic element conversion contained in the (B) ingredient, the (C) ingredient 0.01 - 20 weight sections to (A) ingredient 100 weight section as a It is related with a hardenability constituent given in said either containing the (B) ingredient of desirable embodiment.

[Embodiment of the Invention]

Hereafter, this invention is explained in detail.

Restriction in particular does not have a principal chain skeleton of an organic polymer which has a reactive silicon group used for this invention, and it can use a thing with various kinds of principal ohain skeletons.

condensation polymerization of epsilon-aminoundecanoic acid, Condensation polymerization is carried Polysulfide system polymer; Nylon 610 by the condensation polymerization of the nylon 6 by the ring out from polyamide system polymer,, for example, bisphenol A, and the carbonyl chlorides which have opening polymerization of epsilon caprolactam, hexamethylenediamine, Nyton 66 by the condensation polyisoprene, isoprene or butadiene, acrylonitrile, styrene, etc., Hydrocarbon system polymers, such polyolefine system polymers; Condensation with dibasic acid, such as adipic acid, and glycol, Or the as a hydrogenation polyolefine system polymer produced by hydrogenating polybutadienes or these a two or more-ingredient ingredient among Nylon 12 by the ring opening polymerization of epsilonester produced by carrying out the radical polymerization of the monomers, such as ethyl acrylate polyester system polymer obtained by the ring opening polymerization of lactone; The polyacrylic amino RAURO lactam, and the above-mentioned nylon, such as copolyamide. The polycarbonate and butyl acrylate, Vinyl-base polymers, such as an acrylic ester system copolymer with acrylic ester, such as ethyl acrylate and butyl acrylate, vinyl acetate, aorylonitrile, methyl methacrylate, styrene, etc.; A vinyl monomer is polymerized to said organic polymer. Graft polymer; obtained polyoxypropylene polyoxy butylene copolymer; An ethylene-propylene system copolymer, The Specifically A polyoxyethylene, polyoxypropylene, polyoxy butylene, Polyoxy tetramethylen, a polyoxyethylene polyoxypropylene copolymer, Or polyoxyalkylene series polymers, such as a copolymer of polyisobutylene, isobutylene, isoprene, etc., polychloroprene, A copolymer with polymerization of adipic acid and hexamethylenediamine, and sebacic acid, Nylon 11 by the system polymer manufactured; a diallyl phthalate system polymer etc. are illustrated.

chain skeleton, a polyoxyalkylene series polymer, a hydrocarbon system polymer, a polyester system polymer, a vinyl system copolymer, a polycarbonate system polymer, etc. are preferred. A saturated hydrocarbon system polymer, and a polyoxyalkylene series polymer and a vinyl system copolymer Since acquisition and manufacture are easy among polymers with the above-mentioned principal have a comparatively low glass transition temperature, and their hardened material obtained is especially preferred from excelling in cold resistance.

saturated hydrocarbon system polymers, such as polyisobutylene, hydrogenation polybutadiene, and invention, the reactive silicon group content saturated hydrocarbon system polymer derived from As a saturated hydrocarbon system polymer which has a reactive silicon group used for this hydrogenation polyisoprene, is raised.

basis which can construct a bridge by formation of the siloxane bond which is a reaction which has The reactive silicon group contained in the organic polymer which has a reactive silicon group is a the hydroxyl group or hydrolytic basis combined with the silicon atom, and is accelerated with the http://www4.ipdl.inpit.go.jp/ogi-bin/tran_web_ogi_ejje?atw_u=http%3A%2F%2Fwww4.ipdl.i.. 2010/05/06

parboxylic acid metal salt which is the (B) ingredient.

As a reactive silicon group, it is a general formula (1).

[0031]

[Formula 3]

numbers 1–20. It is the Tori ORGANO siloxy group shown by aryl group of the carbon numbers 6–20, the inside of a formula, R^1 , and R^2 — respectively — independent — an alkyl group of the carbon an aralkyl group of the carbon numbers 7–20, or $\langle R' \rangle_3 SiO-\langle R'$ is substitution or an unsubstituted

ydrocarbon group of the carbon numbers 1~20 independently, respectively). X is a hydroxyf group or a hydrolytic basis independently, respectively. a is 0, 1, 2, or 3, b is 0, 1, or 2, and a and b are not simultaneously set to 0. m — an integer of 0, or 1–19 — it is — a basis expressed is raised.

aminooxy group, a suifhydryl group, an alkenyloxy group, etc. are generally used concrete, for example an acyloxy group, a KETOKISHI mate group, an amino group, an amide group, an acid-amide group, an especially as a hydrolytic basis. A basis for which a hydrogen atom, a halogen atom, an alkoxy group, is not limited but what is necessary is just a conventionally publicly known hydrolytic basis s raised. Among these, although an alkoxy group, an amide group, and an aminooxy group are preferred, a point of hydrolysis nature being quiet and being easy to deal with it to especially an alkoxy group is

preferred

A hydrolytic basis and a hydroxyl group can be combined with one silicon atom in the 1-3 ranges, and a+sigmab) has 1-5 preferred ranges. When a hydrolytic basis and a hydroxyl group join together in two or more] a reactive silicon group, they may be the same and may differ.

Although a silicon atom which forms a reactive silicon group is one or more pieces, in the case of a silicon atom connected by siloxane bond etc., it is preferred that they are 20 or less pieces.

in particular, it is a general formula (13).

Formula 4

 $(\mathbb{R}^2$ and X are the same as the above among a formula.) c is an integer of 1–3. Since the reactive silicon group expressed is easy to receive, it is preferred.

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ethoxy basis, a propoxy group, and an isopropoxy group, is mentioned for X as an example in case c is aralkyl groups, such as aryl groups, such as cycloalkyl groups, such as alkyl groups, such as a methyl siloxy group R' is indicated to be by $_3 {
m SiO}$ – which is a methyl group, a phenyl group, etc. (R'), etc. are group and an ethyl group, and a cyclohexyl group, and a phenyl group, and benzyl, the Tori ORGANO mentioned. In these, the point that the activity of a hydrolysis reaction is high to especially a methyl 3. In the above-mentioned general formula (13), as an example of ${\sf R}^2$ in case ${\sf c}$ is not 3, For example, In the above-mentioned general formula (13), Tori alkoxy silyl groups, such as a methoxy group, an group is preferred.

trimethoxysilyl group can reduce quantity of carboxylic acid metal salt which is the (B) ingredient, it is As more concrete illustration of a reactive silicon group, a trimethoxysilyl group, a triethoxy silyl group, a triisopropoxy silyl group, a dimethoxymethyl silyl group, a diethoxymethylsilyl group, and a diisopropoxy methyl silyl group are mentioned. Its activity is high, and since especially the

What is necessary is just to perform introduction of a reactive silicon group by a publicly known method. That is, the following methods are mentioned, for example.

group content epoxy compound. Subsequently, hydrosilane which has a reactive silicon group is made reactivity to an organic polymer which has functional groups, such as a hydroxyl group, in a molecule to this functional group react, and obtain an organic polymer containing an unsaturation group. Or an unsaturation group content organicity polymer is obtained by copolymerization with an unsaturation (b) Make an organic compound which has an active group and an unsaturation group which show to act on an acquired resultant, and it hydrosilylates.

(**) Make a compound which has a sulfhydryl group and a reactive silicon group react to an organic polymer containing an unsaturation group produced by making it be the same as that of the (b) method

reaotivity to an organic polymer which has functional groups, such as a hydroxyl group, an epoxy (**) Make a compound which has a functional group and a reactive silicon group which show group, and an isocyanate group, in a molecule to this functional group react. A method of making a compound which has a polymer, an isocyanata group, and a reactive silicon group which have a hydroxyl group react to an end a method of (b) or among (**)s from points, like a manufacturing cost becomes low in the above method is preferred. (b) As an example of a hydrosilane compound used in a method, For example, trichlorosilane, methyldi chlorosilicane, dimethylchlorosilicane, Halogenation Silang like phenyl dichlorosilane; Trimethoxysilane, KETOKISHI mate)methylsilane are raised, it is not limited to these. Among these, halogenation Silang and alkoxysilane are especially preferred from points, like availability and hydrolysis reaction nature dimethoxysilane; Methyldi acetoxysilane, The acyloxy silanes like a phenyldiacetoxysilane; although the KETOKISHI mate silanes like bis(dimethyl KETOKISHI mate)methylsilane and bis(cyclohexyl <u> Friethoxysilane, methyldiethoxysilane, methyl dimethoxysilane, The alkoxysilane like phenyl</u> are high.

example, etc. are mentioned, it is not limited in particular. As an example of a compound of having propyltrimethoxysilane, gamma-mercaptpropylmethyl dimethoxysilane, gamma-mercaptopropyl triethoxysilane, gamma-mercapto propylmethyl diethoxysilane, etc. are raised, it is not limited to compound which has a sulfhydryl group and a reactive silicon group as a synthetic method by a radical addition reaction under a radical initiator and/or radical source-of-release existence, for (**) Although a method of introducing into an unsaturation binding site of an organic polymer a said sulfhydryl group and a reactive silicon group, For example, although gamma-mercapto

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propyltrimethoxysilane. For this reason, when three hydrolytic bases, such as a trimethoxysilyl group, As for a silane compound which three hydrolytic bases have combined with one silicon atoms, such as trimethoxysilane, disproportionation may advance. If disproportionation progresses, a remarkable dangerous compound [like] which is dimethoxysilane will arise. However, such disproportionation use a basis combined with one silicon atom as a silicon content group, it is preferred to use a advances in neither gamma-mercapto propyltrimethoxysilane nor gamma-isocyanate synthetic method of (**) or (**).

conversion in GPC (gel permeation chromatography), it is preferred that it is 500 to about 50,000, and about 1,000 to 30,000 liquefied thing which is, carries out and has mobility is especially preferred from exceeded when a number average molecular weight is less than 500, since it is lacking in mobility and points, such as the ease of dealing with it. If sufficient rubber elasticity is not obtained and 50,000 is (A) A number average molecular weight of an organic polymer which is an ingredient, In polystyrene handling is difficult for it, at ordinary temperature, it is not desirable.

not no longer be obtained, if the number of reactive silicon groups contained in a molecule will be less more pieces, and it is preferred that there are 1.1-5 pieces. Since it will become weak firmly and will become scaroe at rubber elasticity if hardenability becomes insufficient, good rubber elasticity may (A) A reactive silicon group in one molecule of organic polymers which are an ingredient is one or than one piece, and five pieces are exceeded, it is not desirable.

may be in both. Since effective network chain density of an organic polymer component contained in a hardened material formed eventually increases especially when a reactive silicon group is in a main A reactive silicon group may be in a main chain terminal or a side chain of an organic polymer, and chain terminal, it is desirable from points, like a rubber-like hardened material of high elongation becomes is easy to be obtained with high intensity.

Said polyoxyalkylene series polymer is a general formula intrinsically (14). :

Formula 5]

alkylene group of the carbon numbers 1-14.) -- it being a polymer which has a repeating unit shown, shape or branched state alkylene group of 2-4. As the example of the repeating unit shown by a among a formula, R³ is a divalent organic group and is the straight chain shape or the branching and, R3 in a general formula (14) has the carbon numbers 1-14, and also preferred straight chain general formula (14),

[0054]

Formula 6]

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JP,4150220,B [DETAILED DESCRIPTION]

** is mentioned. The principal chain skeleton of a polyoxyalkylene series polymer may consist of one kind of repeating unit, and may consist of two or more kinds of repeating units. When used especially for sealant etc., it is desirable from that what comprises the polymer which uses a polyoxypropylene system polymer as the main ingredients is amorphous, or the point which is hypoviscosity

item, a U.S. Pat. No. 3278458 item, A U.S. Pat. No. 3278459 item, a U.S. Pat. No. 3427256 item, a U.S. As a synthetic method of a polyoxyalkylene series polymer, For example, the polymerizing method by shown in a U.S. Pat. No. 3427335 item etc., **********, such as the polymerizing method using a which are shown in JP,61-215623,A react, JP,46-27250,B, JP,59-15336,B, a U.S. Pat. No. 3278457 complex catalyst like a complex produced by making an organoaluminium compound and porphyrin Pat. No. 3427334 item, The polymerizing method by a composite metal cyanide complex catalyst polymerizing method using a catalyst which consists of a phosphazene compound illustrated by an alkali catalyst like KOH, the polymerizing method by a transition metal compound-porphyrin catalyst which consists of a polyphosphazene salt illustrated by JP,H10-273512,A, and the JP,H11-060722,A, are not limited in particular.

ingredients, such as a urethane bond ingredient, may be included in the range which does not spoil an In a principal chain skeleton of the above-mentioned polyoxyalkylene series polymer, other effect of this invention greatly.

and polyol which has a repeating unit of the above-mentioned general formula (14) can be mentioned. tolylene diisocyanate, Aromatic system polyisocyanates, such as diphenylmethane diisocyanate and compounds, such as aliphatic series system polyisocyanates, such as hexamethylene di-isocyanate, xylylene diisocyanate; Isophorone diisocyanate, What is obtained from a reaction of polyisocyanate It is not limited especially as the above-mentioned urethane bond ingredient, but For example,

JP,61-197631,A, JP,61-215622,A, JP,61-215623,A, JP,61-218632,A, JP,H3-72527,A, JP,H3-47825,A, JP.57-164123.A, JP.9-2450.B, a U.S. Pat. No. 3632557 item, What is proposed by each gazette, such as a U.S. Pat. No. 4345053 item, a U.S. Pat. No. 4366307 item, and a U.S. Pat. No. 4960844 item, A manufacturing method of a polyoxyalkylene series polymer which has a reactive silicon group. JP,45–36319,B, JP,46–12154,B, JP,50–156599,A, JP,54–6096,A, JP,55–13767,A, JP,55–13468.A, molecular weights and Mw/Mn (ratio of weight average molecular weight and a number average molecular weight) can use conveniently a polyoxyalkylene series polymer with narrow molecular weight distribution in the 1.6 or less amount of Polymer Division, It is not limited to in particular Although it is indicated by each gazette of JP,H8-231707,A and 6,000 or more number average

A polyoxyalkylene series polymer which has the above-mentioned reactive silicon group may be used alone, and may be used together two or more sorts. [0029]

A polymer which makes a skeleton of a saturated hydrocarbon system polymer which has a reactive

silicon group used for this invention, (1). [whether an olefinic compound of the carbon numbers 1–6, such as ethylene, propylene, 1-butene, and isobutylene, is polymerized as the main ingredients, and] (2) After making diene series, such as butadiene and isoprene, homopolymerize or carrying out copolymerization to the above-mentioned olefinic compound, can obtain by a method of hydrogenating, but. Since an isobutylene system polymer and a hydrogenation polybutadiene system polymer tend to introduce a functional group into an end, and tend to control a molecular weight and can increase the number of end functional groups, they are preferred.

[0061]

All the monomeric units may be formed from an isobutylene unit, and an isobutylene system polymer, It is [in an isobutylene system polymer] desirable, and a monomeric unit which has isobutylene and copolymeric may be especially contained in 10 or less % of the weight of the range preferably 30 or less % of the weight still more preferably 50 or less % of the weight in an isobutylene system polymer. When content of a monomeric unit which has isobutylene and copolymeric exceeds 50 % of the weight, since characteristic physical properties, such as high weatherability, high heat resistance, low moisture permeability, etc. resulting from an isobutylene skeleton, are not fully revealed, it is not desirable.

0062

As such a monomer component, an olefin of the carbon numbers 4–12, vinyl ether, an aromatic vinyl compound, vinylsilane, and arylsilane are raised, for example. As such a copolymer component, for example 1-butene, 2-butene, A 2-methyl-1-butene, a 3-methyl-1-butene, a pentene, 4-methyl-1-pentene, A hexene, a vinylcyclohexene, the methyl vinyl ether, ethyl viryl ether, 1sobutylvinyl ether, styrane, alpha-methylstyrene, dimethylstyrene, Monochlorostyrene, dichlorostyrene, beta-pinene, indene, vinyl trichlorosilane, Vinyl methylsilane, indone, vinyldimethylmethoxysilane, Vinyl rimethylsilane, qivinyl dichlorosilane, divinyl dimethoxysilane, dietavinyl silane, allyltichlorosilane, Allyl methyldi chlorosilicane, allyldimethyl methoxysilane, allyltimethylsilane, allyldimethylsilane, allyldimethylsilane, allyldimethylsilane, diaryl dimethoxysilane, diaryl dimethylsilane, diaryl dimethylsilane, diaryl dimethylsilane, diaryl dimethylsilane, diaryl dimethoxysilane, diaryl dimethylsilane, diaryl dimethoxysilane etc. are raised.

As a monomeric unit which has isobutylene and copolymeric among the above, if vinylsilane and anylsilane are used, a basis which silicon content increases and can act as a silane coupling agent will increase, and the adhesive property of a constituent obtained will improve. [0064]

Other monomeric units may be made to contain besides a monomeric unit used as the main ingredients like a case of the above-mentioned isobutylene system polymer in a hydrogenation polybutadiene system polymer or other saturated hydrocarbon system polymers.

065]

To a saturated hydrocarbon system polymer which has a reactive silicon group used for this invention. A little monomeric units in which an after-polymerization double bond like polyene compounds, such as butadiene and isoprene, remains in the range in which the purpose of this invention is attained may be made to contain in 1 or less % of the weight of the range preferably especially 5 or less % of the weight still more preferably 10 or less % of the weight.

A saturated hydrocarbon system polymer which has these reactive silicon groups can be independent, or can be used together two or more sorts.

[0067]

A process of a saturated hydrocarbon system polymer which has a reactive silicon group next is explained.

[0068]

An isobutylene system polymer which has a reactive silicon group in molecular chain terminals among sobutylene system polymers which have a reactive silicon group, an end organic-functions type obtained by the polymerizing method (cationio polymerization method using a specific compound combining and [which is called iniphor] and a chain transfer agent) called iniphor method — it can manufacture preferably using a whole end organic-functions type isobutylene system polymer. As a

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process of a saturated hydrocarbon system polymer which has a reactive silicon group, For example, general formula (15) after obtaining polyisobutylene which has an unsaturation group at the end by dehydrohalogenation reaction of an end of a polymer which has third class carbon-chlorine combination obtained by a polymerization reaction, the reaction of an end of a polymer and allyi trimethylsilanc which have third class carbon-chlorine combination, etc.:

[0069] [Formula 7]

$$H \leftarrow \text{Si-O} - \text{Si-X}_{\text{a}} \qquad (15)$$

(R¹, R², X, m, a, and b are the same as the above among a formula.) — the hydrosilane compound (this compound is a compound which the hydrogen atom combined with the basis expressed with a general formula (1).) expressed — desirable — general formula (16):

[Formula 8]

(16)

 R^2 , X, and c are the same as the above among a formula.) — it can obtain by the reaction (hydrosilylation reaction) to which the hydrosilane compound expressed is made to add using a platinum catalyst.

[007

As a hydrosilane compound, for example Trichlorosilane, methyldi chlorosilicane, Halogenation Silang like dimethylchilorosilicane and phenyl dichlorosilane; Trimethoxysilane, Triethoxysilane, methyldischoxysilane, methyl dimethoxysilane, The alkoxysilane like phenyl dimethoxysilane; Methyldi acetoxysilane, The acyloxy silanes like a phenyldiacetoxysilane; although the KETOKISHI mate silanes like bis(dimethyl KETOKISHI mate)methylsilane and bis(cyclohexyl KETOKISHI mate)methylsilane are raised, it is not limited to these. Among these, halogenation Silang and alkoxysilane are preferred especially in respect of the ease of acquisition, etc.

Louzel Such a manufacturing method is indicated in each gazette of JP,4–69659,B, JP,7–108928,B, JP,63– 254149,A, JP,64–22904,A, and the patent No. 2539445, for example. Lan isobutylene system polymer which has a reactive silicon group in a chain side chain is manufactured by adding and carrying out copolymerization of the vinylsilane and arylsilane which have a reactive silicon group into a monomer containing isobutylene.

In the case of a polymerization reaction which manufactures an isobutylene system polymer which has a reactive silicon group in molecular chain terminals. After carrying out copolymerization of the vinylsilane and arylsilane etc. which have a reactive silicon group in addition to an isobutylene monomer which is the main ingredients, an isobutylene system polymer which has a reactive silicon group in an end and a chain side chain is manufactured by introducing a reactive silicon group into an

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As the vinylsilane which has a reactive silicon group, and arylsilane, For example, vinyl trichlorosilane, allyldimethylchlorosilicane, Allyldimethyl methoxysilane, diaryl dichlorosilane, diaryl dimethoxysilane, vinyl methyldi chlorosilicane, vinyldimethylchlorosilicane, Vinyldimethylmethoxysilane, divinyl gamma-methacryloyl oxypropyl trimethoxysilane, gamma-methacryloyl oxypropyl methyl dichlorosilane, divinyl dimethoxysilane, Allyltrichlorosilane, allyl methyldi chlorosilicane, dimethoxysilane, etc. are raised.

has an end olefin group uses a hydroxyl group of an end hydroxy hydrogenation polybutadiene system polymer as oxy metal groups, such as -ONa and -O.K., first, for example: group. A hydrogenation polybutadiene polymerization object which has a reactive silicon group can be acquired by the hydrosilylation reaction of a hydrogenation polybutadiene polymerization object which has an olefin group. General formula (17) after a hydrogenation polybutadiene system polymer which in this invention, a hydrogenation polybutadiene polymerization object which has a reactive silicon group can be mentioned as a saturated hydrocarbon system polymer which has a reactive silicon

CH₂=CH-R⁴-Y (17)

and Y is "R5-, -R5-000-, or -R5-00. - (R5 is a divalent hydrocarbon group of the carbon numbers Halogen atoms, such as a chlorine atom, a bromine atom, and iodine atoms, and R^4 are among [type, I-20, and) an alkylene group, a cyclo alkylene group, an allylene group, and an aralkylene group --- it is desirable --- by a divalent organio group shown. -- It can obtain by making an organic halogenated compound shown by especially a divatent basis chosen from GH $_2$ - and -R $^{\circ}$ -C $_6$ H $_4$ -CH $_2$ - (R $^{\circ}$ is a hydrocarbon group of the carbon numbers 1-10) being preferred] react.

system polymer as a oxy metal group, A method of making it react to alkaline-water oxides, suoh as metal alkoxide;NaOH like metal hydride;NaOCH₃ like Na and alkaline metal;NaH like K and KOH, etc. is As a method of using terminal hydroxyl groups of an end hydroxy hydrogenation polybutadiene

raised.

a general formula (17) after that, A hydrogenation polybutadiene system polymer which is the amount weight as an end hydroxy hydrogenation polybutadiene system polymer used as a starting material is organic halogenated compound which contains two or more halogen in one molecule, and is shown by Although an end olefin hydrogenation polybutadiene system polymer with the almost same molecular react to obtain a polymer of the amount of Polymer Division more, at the time, such as a methylene nalogenated compound which can increase a molecular weight if it is made to react to a multivalent obtained in said method, Before making an organic halogenated compound of a general formula (17) chloride, bis(chloromethyl)benzene, and bis(chloromethyl)ether. If it is made to react to an organic of Polymer Division more, and has an olefin group at the end can be obtained.

As an example of an organic halogenated compound shown by said general formula (17), For example, (chloromethyl) ether, 1-hexenyl(chloromethoxy) benzene, allyloxy (chloromethyl) benzene, etc. are raised, it is not limited to them. Among these, an allyl chloride is cheap, and since it moreover reacts an allyl chloride, an allyl star's picture, vinyl(chloromethyl) benzene, Allyl(chloromethyl) benzene, allyl bromomethyl) benzene, Although allyl (chloromethyl) ether, allyl(chloromethoxy) benzene, 1-butenyl easily, it is desirable.

polymer may be manufactured by carrying out an addition reaction using a platinum system catalyst in a hydrosilane compound like a case of an isobutylene system polymer which has a reactive silicon introduction of a reactive silicon group to said end olefin hydrogenation polybutadiene system group in molecular chain terminals.

above does not contain substantially an unsaturated bond which is not an aromatic ring in a molecule, Compared with a sealing agent etc. which consist of a conventional rubber system polymer like an When a saturated hydrocarbon system polymer which has a reactive silicon group as mentioned

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weatherability becomes good remarkably. Since this polymer is a hydrocarbon system polymer, its water resisting property is good, and a low hardened material of humidity permeability is obtained. organic system polymer or an oxyalkylene system polymer which has an unsaturated bond,

together two or more sorts. Specifically, a polyoxyalkylene series polymer which has a reactive silicon polymer which has a reactive silicon group, and an organic polymer which blends two or more sorts An organio polymer which has these reactive silicon groups may be used alone, and may be used group, a saturated hydrocarbon system polymer which has a reactive silicon group, a vinyl-base chosen from a group, ** and others, can also be used.

A manufacturing method of an organic polymer which blends a polyoxyalkylene series polymer which has a reactive silicon group, and a vinyl-base polymer which has a reactive silicon group, Although indicated by JP,59-122541,A, JP,63-112642,A, JP,H6-172631,A, JP,H11-116763,A, etc., it is not limited to in particular these.

A desirable example has a reactive silicon group and a chain is a following general formula substantially (18)::

[Formula 9]

$$\begin{array}{ccc} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ \end{array}$$

(a hydrogen atom or a methyl group, and ${
m R}^7$ show the alkyl group of the carbon numbers 1-8 among a The acrylic ester monomer unit which has an alkyl group of the carbon numbers 1–8 expressed with formula, as for R^6) (meta), and following general formula (19) :

[Formula 10]

(-- inside of formula, and R^6 — the above — the same -- R^3 shows a with a carbon numbers of ten or more alkyl group —) — to the copolymer which consists of an acrylic ester monomer unit which blending and manufacturing the polyoxyalkylene series polymer which has a reactive silicon group. has a with a carbon numbers of ten or more expressed alkyl group (meta-). It is the method of Acrylic acid (meta) expresses acrylic acid and/or methacrylic acid with the above-mentioned expressive form.

propyl group, n-butyl group, t-butyl group, a 2-ethylhexyl group, etc. --- desirable --- 1-4 --- an alkyl as ${
m R}^7$ of said general formula (18) — the carbon numbers 1–8 of a methyl group, an ethyl group, a group of 1–2 is raised still more preferably. An alkyl group of \mathbb{R}^7 may be independent and may be mixed two or more sorts. as R^8 of said general formula (19) — ten or more carbon numbers of a lauryl group, a tridecyl group, a

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setyl group, a stearyl group, a behenyl group, etc. — usually --- 10-30 --- a long-chain alkyl group of 0-20 is raised preferably. Like a case of $\rm R^7$, an alkyl group of $\rm R^8$ may be independent and may be mixed two or more sorts.

formula (19) substantially, a "real target" here means that the sum total of a monomeric unit of a formula (18) which exists in this copolymer, and a formula (19) surpasses 50 % of the weight. The sum total of a monomeric unit of a formula (18) and a formula (19) is 70 % of the weight or more Although a chain of this vinyl system copolymer consists of a monomeric unit of a formula (18) and a preferably.

From a point of compatibility with a polyoxyalkylene series polymer, as for an abundance ratio of a monomeric unit of a formula (19), 95:5-40:60 are preferred at a weight ratio, and 90:10–60:40 are still more preferred.

[1600]

methylstyrene, alkyl vinyl ether, VCM/PVC, vinyl acetate, vinyl propionate, ethylene, etc. in addition Diethylamino ethyl acrylate, diethylamino ethyl methacrylate, A monomer containing amino groups, such as aminoethyl vinyl ether, a monomerio unit which originates in acrylonitrile, styrene, alpha-As monomeric units other than a formula (18) which may be contained in this copolymer, and a formula (19), For example, carboxylic acid groups, such as acrylic acid and methacrylic acid, methylolmethacrylamide, Epoxy groups, such as glycidyl acrylate and glycidyl methacrylate, acrylamide, Amide groups, such as methacrylamide, N-methylolacrylamide, and Nto this is raised.

has a reactive silicon group, and a vinyl-base polymer which has a reactive silicon group is indicated Although an organic polymer which blends a saturated hydrocarbon system system polymer which by JP H1-168764,A, JP,2000-186176,A, etc., it is not limited to in particular these.

organic polymer which has a reactive silicon group as a manufacturing method of an organic polymer which blends a vinyl-base polymer which has a reactive silicon group can be used. Although this A method of polymerizing an acrylic ester (meta) system monomer elsewhere under existence of an manufacturing method is concretely indicated by each gazette, such as JP,59–78223,A, JP,59– 68014,A, JP,60-228516,A, and JP,60-228517,A, it is not limited to these.

manganese, carboxylio acid nickel, carboxylic acid cobalt, and a carboxylic acid zirconium, It functions as what is called a silanol condensation catalyst in which a siloxane bond may be made to form from One or more sorts of carboxylic acid metal salt chosen from carboxylic acid barium, carboxylic acid Carboxylio acid vanadium, carboxylic acid iron, carboxylic acid titanium, carboxylic acid potassium, Carboxylic acid calcium used as a (B) ingredient in a hardenability constituent of this invention, a hydroxyl group combined with a silicon atom contained in an organio polymer which is the (A) ngredient of this invention, or a hydrolytic basis.

catalyst is high, carboxylic acid calcium, carboxylic acid vanadium, carboxylic acid iron, carboxylic acid in said carboxylic acid metal salt, carboxylic acid calcium, carboxylic acid vanadium, Carboxylic acid manganese, and a carboxylic acid zirconium, it is more desirable from a point that the activity of a ron, oarboxylic acid titanium, carboxylic acid potassium, carboxylic acid barium, carboxylic acid titanium, and a carboxylic acid zirconium are still more preferred, and carboxylic acid iron and carboxylic acid titanium are especially the most preferred.

potassium, carboxylic acid barium, carboxylic acid manganese, carboxylic acid nickel, carboxylic acid nardenability constituent obtained, and a point that the heat resistance of a hardened material and weatherability which are obtained are high, and carboxylic acid calcium, carboxylic acid titanium, carboxylic acid potassium, carboxylic acid barium, and a carboxylic acid zirconium are still more cobalt, and a carboxylic acid zirconium, it is more desirable from a point with little coloring of a Carboxylic acid calcium, carboxylic acid vanadium, carboxylic acid titanium, Carboxylic acid

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JP,4150220,B [DETAILED DESCRIPTION]

This (B) ingredient uses as the main ingredients carboxylic acid metal salt expressed with gencral formula (2) - (12), respectively. Ca(OCOR) 2 (2)

V(OCOR) 3 (3)

Fe(OCOR) 2 (4) Fe(OCOR) 3 (5)

Ti(OCOR) 4 (6)

K(000R) (7)

Ba(OCOR) 2 (8)

Mn(OCOR), (9)

nickel(OCOR) , (10)

Zr (O) (OCOR), (12) Co(OCOR) 2 (11)

(The inside R of a formula is substitution or an unsubstituted hydrocarbon group, and may include a carbon carbon double bond.)

As carboxylic acid, a carboxylic acid group content compound of a hydrocarbon system of 2-40 is carboxylic acid group content compound of a hydrocarbon system of the carbon numbers 2-20 used suitably, and a carbon number including carbonyl carbons may be especially used for a suitably from a point of availability here.

tuberculostearic acid, a pivalic acid, and neo decanoic acid; A tariric acid, Fatty acid with triple bonds, Enanthic acid, caprylic acid, 2-ethylhexanoic acid, pelargonic acid, capric acid, Undecanoic acid, lauric 10,12-octadecadienoic acid, HIRAGO acid, alpha-eleostearic acid, beta-eleostearic acid, punicic acid, hydroxyhexadecanoic acid, YARAPI Norian acid, uni-PERIN acid, AMBURETTORU acid, ARYURITTO linolenic acid, 8,11,14-eicosatriencic acid, a 7,10,13-docosatriencic acid, 4,8,11,14-hexadeca tetraencic acid, MOROKUCHI acid, steer RIDON acid, Arachidonic acid, 8,12,16,19-docosatetraencic acid, 4,8,12,15,18-eicosapentaenoic acid, Polyene unsaturated fatty acid, such as clupanodonic acid, Oxygenated fatty acid, such as 9,10-dihydroxyootadecanoic acid, recinoleic acid, cam ROREN acid, acid, Tsuzuic acid, FIZETERIN acid, myristoleic acid, 2-hexadecenoic acid, 6-hexadecenoic acid, 7-Naphthenic acid, A malvalic acid, sterculic acid, HIDONO carbyne acid, chaulmoogric acid, Alicyclic When it illustrates concretely, acetio acid, propionic acid, butanoic acid, a valeric acid, caproic acid, vaccenic acid, gadoleic acid, Gondo Inn acid, a cetoleic acid, erucic acid, brassidio acid, selacholeic acid, Straight chain saturated fatty acid groups, suoh as RAKUSERU acid; Undecylenic acid, Linder nonadecanoic acid, arachin acid, Behenic acid, lignoceric acid, cerinic acid, montanic acid, melissic licanic acid, ferron acid, and cerebronic acid; dicarboxylic acid, such as succinic acid, glutaric acid, hexadecenoio acid, palmitoleic acid, a petroselinic acid, Oleic acid, elaidic acid, ASUKUREPIN acid, carvone acids, such as gorlic acid; Sabinic acid, 2-hydroxytetradecanoic acid, IPURORU acid, 2acid, Monoene unsaturated fatty acid, such as KISHIMEN acid and RUMEKUEN acid; Linolic acid, acid, 2-hydroxyoctadecanoic acid, 12-hydroxyoctadecanoic acid, 18-hydroxyoctadecanoic acid, such as steer roll acid, a crepenynic acid, KISHIMENIN acid, and 7-hexa crepe-de-Chine acid; acid, tridecylacid, myristic acid, pentadecyl acid, Pulmitic acid, heptadecylacid, stearic acid, herring acid, and docosahexaenoic acid; Iso acid, Branch fatty acid, such as anteiso acid, adipic acid, pimelic acid, SUPERIN acid, azelaic acid, and sebacic acid, is mentioned.

radical (workability --- bad). Therefore, as for the melting point of said carboxylic acid, it is preferred that it is 65 ** or less, it is more preferred that it is -50-50 **, and it is preferred that it is especially when the melting point of said carboxylic acid is high (crystallinity is high), the melting point becomes high in a similar manner, and it is hard to deal with carboxylic acid metal salt which has the acid

when a carbon number of said carboxylic acid is large (a molecular weight is large), it becomes

solid state or viscosity and which is hard to deal with it (workability -- bad). On the contrary, when a extended a constituent thinly, volatilization by heating may be large and catalyst ability of carboxylic numbers including carbon of a carbonyl group are 2-17, it is more preferred that it is 3-13, and it is iquefied and carboxylic acid metal salt which has the acid radical becomes a thing which has a high carboxylic acid metal salt may fall [carboxylic acid metal salt which has the acid radical], including mostly an ingredient which volatilizes easily with heating. Especially on conditions (thin layer) which acid metal salt may fall greatly. Therefore, as for said carboxylic acid, it is preferred that carbon carbon number of said carboxylic acid is small (a molecular weight is small), catalyst ability of preferred that it is especially 5-10.

good to said carboxylic acid has 2-ethylhexanoic acid, octylic acid, neo decanoic acid, oleic acid, or Especially acquisition is easy, and is cheap and a point that compatibility with the (A) ingredient is preferred naphthenic acid. Said naphthenic acid is expressed with an empirical formula (20).

carboxylic acid (2-ethylhexanoic acid etc.) whose carbon atoms in which said carboxylic acid adjoins quaternary carbon A pivalic acid etc. are more preferred from a cure rate being quick, and especially carboxylic acid whose carbon atom which adjoins a carbonyl group is quaternary carbon is preferred. a carbonyl group are the third class carbon and carboxylic acid (neo decanoic acid.) which is [0102]

(divalent), naphthenic acid nickel (divalent), cobalt naphthenate (divalent), a naphthenic acid zirconium cobalt (divalent), 2-ethylhexanoic acid zirconium (tetravalence), neo decanoic acid iron (divalent), Neo ethylhexanoic acid manganese (divalent), 2-ethylhexanoic acid nickel (divalent), 2-ethylhexanoic acid decanoic acid iron (trivalent), neo decanoic acid titanium (tetravalence), neo decanoic acid vanadium (divalent), oleic acid iron (trivalent), oleic acid titanium (tetravalence), Oleic acid vanadium (trivalent), oleic acid calcium (divalent), oleic acid potassium (univalent), oleic acid barium (divalent), manganese salt, 2-ethylhexanoio acid iron (divalent), 2-ethylhexanoic acid iron (trivalent), 2-ethylhexanoic acid Naphthenic acid potassium (univalent), naphthenic acid barium (divalent), manganese naphthenate From a viewpoint of availability and compatibility, as an example of desirable carboxylic acid metal (divalent), 2-ethylhexanoic acid potassium (univalent), 2-ethylhexanoic acid barium (divalent), 2titanium (tetravalence), 2-ethylhexanoic acid vanadium (trivalent), 2-ethylhexanoic acid calcium (tetravalence), naphthenic acid iron (divalent), naphthenic acid iron (trivalent), Naphthenic acid (trivalent), Neo decanoic acid caloium (divalent), neo decanoic acid potassium (univalent), Neo oleate (divalent), oleic acid nickel (divalent), oleic acid cobalt (divalent), An oleic acid zirconium decanoic acid barium (divalent), a neo decanoic acid zirconium (tetravalence), Oleic acid iron titanium (tetravalence), naphthenio acid vanadium (trivalent), caloium naphthenate (divalent), (tetravalence), etc. are mentioned.

2-ethylhexanoic acid iron (trivalent), neo decanoic acid iron (trivalent), oleic acid iron (trivalent), and Naphthenic acid iron (trivalent) and naphthenic acid titanium (tetravalence) are more preferred, and 2-ethylhexanoic acid iron (divalent) from a viewpoint of oatalytic activity, 2-ethylhexanoic acid iron decanoic acid iron (trivalent), neo decanoic acid titanium (tetravalence), oleic acid iron (divalent), trivalent), 2-ethylhexanoic acid titanium (tetravalence), neo decanoic acid iron (divalent), Neo Oleic acid iron (trivalent), oleic acid titanium (tetravalence), naphthenic acid iron (divalent), especially naphthenic acid iron (trivalent) are preferred.

[0104]

calcium (divalent), 2-ethylhexanoic acid potassium (univalent), 2-ethylhexanoic acid barium (divalent), divalent), a neo decanoic acid zirconium (tetravalence), Oleic acid titanium (tetravalence), oleic acid decanoic acid calcium (divalent), Neo decanoic acid potassium (univalent), neo decanoic acid barium from a viewpoint of coloring to 2-ethylhexanoic acid titanium (tetravalence). 2-ethylhexanoic acid zirconium (tetravalence), naphthenic acid titanium (tetravalence), calcium naphthenate (divalent), naphthenic acid potassium (univalent), naphthenic acid barium (divalent), and a naphthenic acid 2-ethylhexanoic acid zirconium (tetravalence), Neo decanoic acid titanium (tetravalence), neo calcium (divalent), oleic acid potassium (univalent), Oleic acid barium (divalent), an oleic acid zirconium (tetravalence) are more preferred.

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JP,4150220,B [DETAILED DESCRIPTION]

method to which a carboxylic acid group content compound and metal powder are made to react,] an this In addition, a sedimentation method which settles metallic soap, A carboxylic acid group content sodium hydroxide react, and makes solution of hard soap, Solution of metal salt prepared apart from compound or its ester and metaled hydroxide, an oxide, A method etc. to which an alcoholate or a Such carboxylic acid metal salt makes a carboxylic acid group content compound, or its ester and soorification to which a weak acid salt is made to react at an elevated temperature, and direct chloride, and a carboxylic acid group content compound are made to react in [else /, such as anhydrous organic solvent are taken. As for such carboxylic acid metal salt, it is preferred to dilute with diluting solvents, such as a mineral spirit, toluane, hexylene glyool, a diethylene glycol, a paraffin oil, and dioctyl phthalate, and to be used with a gestalt of a solution whose metal content is about 1 to 40 % of the weight.

[0107]

workability except that a good hardened material will become generation of heat and foaming local at (B) About 0.005–5 weight sections are preferred at metallic element conversion contained in the (B) ingredient to (A) ingredient 100 weight section as amount of ingredient used, and also about 0.01-3 weight sections are preferred. (B) Since a cure rate may become slow and a hardening reaction will desirable. On the other hand, pot life becomes short too much and is not preferred from a point of the time of hardening arise, and is hard to be obtained, if loadings of the (B) ingredient exceed this become fully difficult to advance if loadings of an ingredient are less than this range, it is not

using it alone, it can be used combining two or more sorts, and also can use together with carboxylic manganese, carboxylic acid nickel, carboxylic acid cobalt, and a carboxylic acid zirconium, Besides acid tin salt, carboxylic acid lead salt, carboxylic acid bismuth salt, carboxylic acid ceric salt, etc. The aforementioned carboxylic acid calcium, carboxylic acid vanadium, carboxylic acid iron, Carboxylic acid titanium, carboxylic acid potassium, carboxylic acid barium, carboxylic acid

(dimethyl aminomethyl) phenol, Although morpholine, N-methylmorpholine, 2-ethyl-4-methylimidazole, example, Specifically Methylamine, ethylamine, propylamine, isopropylamine, A butylamine, amyl amine, On the other hand, only with carboxylic acid metal salt of the (B) ingredient, activity is low, and when hexylamine, octylamine, 2-ethylhexylamine, Nonyl amine, decyl amine, lauryl amine, pentadecyl amine, moderate hardenability is not acquired, various amine compounds which are the (C) ingredients as a dodecamethylenediamine, dimethylethylenediamine, Triethylenediamine, guanidine, diphenylguanidine, co-catalyst can be added. As various amine compounds, although indicated to JP,H5-287187,A, for Dimethylamine, Diethylamine, dipropyl amine, diisopropylamine, dibutyl amine, Diamylamine, dioctyl aromatic-amine; Lauryl aniline, stearylaniline, a triphenylamine, N.N-dimethylaniline, dimethylbenzyl benzylamine, Diethylamino propylamine, xylylene diamine, ethylenediamine, Hexamethylenediamine, N,N,N', and N'-tetramethyl 1,3-butanediamine, N,N,N', N'-tetramethyl ethylene diamine, 2,4,6-tris amine, and trioctylamine, Triaryl amine, Aliphatic series unsaturation amines, such as oleylamine; butylstearylamine; Triethylamine, Aliphatic series tertiary amines, such as triamylamine, trihexyl amine, di(2-ethylhexyl) amine, Didecyl amine, dilauryl amine, JISECHIRU amine, distearyl amine, , and 8-diazabicyclo (5, 4, 0) undecene 7 (DBU) etc. are mentioned, it is not limited to these. Aliphatic series primary amines, such as Sept Iles amine, stearylamine, and cyclohexylamine; dimethylamino ethanol, diethylenetriamine, Triethylenetetramine, tetraethylenepentamine, aniline, etc. reach, As other amines, monoethanolamine, diethanolamine, Triethanolamine, Aliphatic series secondary amines, such as methylstearylamine, ethylstearylamine, and

combined. As an example of this hydrolytic silicon group, X can mention a thing which is a hydrolytic In this invention, an amino group content silane coupling agent can also be used as a (C) ingredient. hydrolytic basis can specifically be mentioned, a methoxy group, an ethoxy basis, etc. are preferred basis among bases expressed with a general formula (1). Although a basis already illustrated as a from a point of a hydrolysis rate. As for especially the two or more number of a hydrolytic basis, Said amino group content silane coupling agent is a compound which has a basis (henceforth a hydrolytic silicon group) and an amino group containing a silicon atom which a hydrolytic basis

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three or more pieces are preferred.

As an example of an amino group content silane coupling agent, gamma-aminopropyl trimethoxysilane, aminopropyl trimethoxysilane, gamma-(2-aminoethyl) aminopropyl methyl dimethoxysilane, gamma-(2vinylbenzyi gamma-aminopropyl triethoxysilane, etc. can be mentioned. Amino modifying silyl polymer which is the denatured derivative, silanizing amino polymer, an unsaturation aminosilane complex, a gamma-aminopropyl triethoxysilane, gamma-aminopropyl triisopropoxy silane, gamma-aminopropyl mentioned amino group content silane coupling agent may be used only by one kind, and may carry gamma-(2-aminoethyl) aminopropyl triisopropoxy silane, gamma-ureido propyltrimethoxysilane, Naminoethyl) aminopropyl triethoxysilane, gamma~(2-aminoethyl) aminopropyl methyldiethoxysilane, phenyl-gamma-aminopropyl trimethoxysilane, N-benzyl-gamma-aminopropyl trimethoxysilane, Nphenylamino long chain alkyl silane, amino silanizing silicone, etc. can use these. The abovemethyl dimethoxysilane, gamma-aminopropyl methyldiethoxysilane, gamma-(2-aminoethyl) out two or more kind mixing use.

As these (C) ingredients, since co-catalyst ability changes greatly with structure of the (C) ingredient system polymer as a (A) ingredient, aliphatic series secondary amines, such as comparatively long suitable according to a kind of (A) ingredient to be used. For example, when using an isobutylene tself, compatibility with the (A) ingredient, etc., it is preferred to choose a compound which was chain aliphatic series secondary amines, such as dioctyl amine and distearyl amine, and dicyclohexylamine, are preferred from a point that co-catalyst ability is high.

weight sections are preferred to organic polymer 100 weight section of the (A) ingredient, and also its 0.1 - 5 weight section is more preferred. A cure rate may become it slow that loadings of an amine advance. On the other hand, if loadings of an amine compound exceed 20 weight sections, pot life As for loadings of an amine compound which is the aforementioned (G) ingredient, about 0.01–20 compound are less than 0.01 weight sections, and a hardening reaction becomes fully difficult to may become short too much and is not preferred from a point of workability.

Silane coupling agents other than an amino group content silane coupling agent can also be used for a constituent of this invention. As functional groups other than an amino group, a sulfhydryl group, an epoxy group, a carboxyl group, a vinyl group, an isocyanate group, isocyanurate, halogen, etc. can be illustrated.

chloropropyltrimetoxysilane,]. --- isocyanurate silanes [, such as tris (trimethoxysilyl) isocyanurate,]. isocyanate propylmethyl dimethoxysilane, can be mentioned. A block isocyanate silane, silanizing polyester, etc. which are the derivatives which denaturalized these can be used as a silane coupling propyltriethoxysilane, gamma-glycidoxy propyl methyldimethoxysilane, beta-(3, 4-epoxycyclohexyl) gamma-mercapto propyltrimethoxysilane, gamma-mercaptopropyl triethoxysilane, Sulfhydryl group oxypropyl methyl dimethoxysilane, vinyl type unsaturation group content Silang [, such as gamma-As an example of silane coupling agents other than an amino group content silane coupling agent, isocyanatepropyl triethoxysilane, gamma-isocyanate propylmethyl diethoxysilane, and gammaisocyanate group content Silang, such as gamma-isocyanate propyltrimethoxysilane, gamma-AKURO yloxy propylmethyl triethoxysilane,]: — containing halogen Silang [, such as gammaaminopropyl trimethoxysilane; Vinyltrimetoxysilane, vinyltriethoxysilane, gamma-methacryloyl content Silang, such as gamma-mercaptpropylmethyl dimethoxysilane and gamma-mercapto methoxyethoxy)Silang, Carboxysilanes, such as N-beta-(carboxymethyl) aminoethyl gammaethyltrimethoxysilane, Epoxy group content Silang, such as beta-(3,4-epoxycyclohexyl) ethyltriethoxysilane; beta-carboxyethyl triethoxysilane, beta-carboxyethyl phenylbis(2propylmethyl diethoxysilane; Gamma-glycidoxypropyltrimetoxysilane, gamma-glycidoxy

The amount of [in case used of using silane coupling agents other than an amino group content silane coupling agent] has 0.01 - 20 preferred weight section to (A) ingredient 100 weight section.

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and also its 0.1 - 5 weight section is more preferred.

carbonate, clay, talo, titanium oxide, aluminium hydroxide, magnesium carbonate, aluminum impalpable titaniun oxide, tale, etc. are more preferred than points, such as a reinforoing effect, the increase-inquantity effect, and an ease of acquisition. These fillers may be used independently and may be used Various fillers can be used for a hardenability constituent of this invention if needed. As an example together two or more sorts. The amount of [in case used of using a filler] has 10-1000 preferred weight section to (A) ingredient 100 weight section, and its 50-300 weight section is still more mica, walnut shell powder, chaff powder, Graphite, diatomite, clay, fume silica, sedimentation nature powder, the Flint powder, the end of zinc dust, etc. are raised. Among these fillers, sedimentation nature silica, fume silioa, orystalline silica, fused silica, dolomite, carbon black, calcium carbonate, of said filler, for example Wood flour, PARUBU, a cotton chip, Asbestos, glass fiber, carbon fiber, silica, Crystalline silica, fused silica, dolomite, a silicic acid anhydride, carbon black, calcium

phthalate; Dioctyl adipate, *** aliphatic dibasic acid ester, such as succinic acid isodecyl and dioctyl sebacate; Diethylene glycol dibenzoate, **** glycol ester, such as pentaerythritol ester; Butyl oleate, bulking agent, it is more effective. As an example of a plasticizer, dioctyl phthalate, dibutyl phthalate, phosphoric ester, such as trioctyl phosphate and phosphoric acid octyldiphenyl; Epoxidized soybean oil, **** epoxy plasticizers, such as epoxidation linseed oil and epoxy stearic acid benzyl; Polyether, such as a polyester plasticizer, polypropylene glycol which is polyester of dibasic acid and dihydric Polybutadiene, Butadiene Acrylonitrile, polychloroprene, polyisoprene, polybutene, hydrogenation **** aliphatic series ester species, such as methyl acetyl ricinolate; Tricresyl phosphate, **** polybutadiene, hydrogenation polyisoprene, Hydrocarbon system oligomer, such as process oil; enlarged or a lot of bulking agents can be mixed if a plasticizer is used, using it together with a In a hardenability constituent of this invention, since elongation of a hardened material can be **** phthalic ester, such as butylbenzyl phthalate, di-isodecyl phthalate, and JIISO undecyl alcohol, and its derivative; [Polly alpha-methylstyrene,] Polystyrene, such as polystyrene; chlorinated paraffins are illustrated.

These plasticizers may be used alone and may be used together two or more sorts. A desirable result will be obtained if the amount of plasticizers is used in the range of one to 200 weight section to reactive-silicon-group-containing-organic-polymer 100 weight section. [0121]

An epoxy resin can be used together in a hardenability constituent of this invention. In this case, an

epoxy resin and reactive silicon group containing organic polymer can be reformed. [0122]

such as tetrahydrophtal acid diglycidyl ester and diglycidyl hexahydrophthalate, m-aminophenol series As an epoxy resin, can use a publicly known thing widely conventionally, and For example, a bisphenol A type epoxy resin, Fire retardancy type cpoxy resins, such as bisphenol F type epoxy resin and epoxy resin, A glyoidyl ether type epoxy resin of a bisphenol A propylene oxide addition, Diglycidyl pglycidyl ether of tetrabromobisphenol A, Novolak type epoxy resin, a hydrogenation bisphenol A type epoxy resin, a diaminodiphenylmethane system epoxy resin, Urethane modified epoxy resin, various alcohol, such as polyalkylene glycol diglycidyl ether and glycerin, a hydantoin type epoxy resin, and oxybenzoic acid, phthalic acid diglycidyl ester, Phthalic acid diglycidyl ester system epoxy resins, isocyanurate, An epoxidation thing of unsaturation polymers, such as glycidyl ether of polyhydric cycloaliphatic-epoxy-resin, N, and N-diglycidyl aniline, N,N-diglycldyl o-toluidine, triglycidyl petroleum resin, etc. can be mentioned. It is desirable from a point of a thing containing at least two epoxy groups having high reactivity when hardening, and a hardened material tending to form the three-dimensional network structure into a bisphenol F type epoxy resin, novolak type epoxy resin, and a phthalic acid diglycidyl ester system molecule, also in these epoxy resins. As a desirable epoxy resin, a bisphenol A type epoxy resin, epoxy resin can be illustrated from availability and an adhesive point.

Diaminodiphenylmethane, diaminodiphenyl sulfone, isophoronediamine, Amines, suoh as 2,4,6-tris (dimethyl aminomethyl) phenol, Tertiary amine salts, polyamide resin, ketimines, and aldimine. Insidious hardening agents, such as enamines, imidazole derivatives, and dicyandiamides. Boron trifluoride complex compounds, phthalic anhydride, hexahydro phthalic anhydride, Anhydrous carboxylic acid, such as tetrahydro phthalic anhydride and methylene tetrahydro phthalic anhydride, and carboxylic acid, such acid can be mentioned.

An epoxy resin receives reactive—silicon–group—containing—organic—polymer 100 weight section, it is preferred to use it in the range of one to 100 weight section and also ten to 50 weight section, and it is preferred to blend a hardening agent of an epoxy resin per epoxy resin 100 weight section in the range of one to 200 weight section and also ten to 100 weight section.

An ingredient which limitation in particular does not have in the method of preparation of a hardenability constituent of this invention, for example, was described above is blended, it kneads under ordinary temperature or heating using a mixer, a roll, a kneader, etc., or an ingredient is dissolved using a little suitable solvents, and a usual method of mixing may be adopted. A 1 liquid type, two-component type, or many liquid type compound can also be built and used by combining these ingredients suitably.

[0127]
If a hardenability constituent of this invention is exposed into the atmosphere, by operation of moisture, it will form network structure in three dimensions, and will harden it to a solid which has reabler—like elasticity.

In a hardenability constituent of this invention, various additive agents can be added if needed. Other curing catalysts as an example of said additive for example (for example, tetravalent tin, divalent tin, etc.), Adhesive grant agents which adjust the tractive characteristics of a hardened material to generate, such as a physical-properties regulator and a silane coupling agent, An antioxidant, radical inhibitor, an ultraviolet ray absorbent, a metal deactivator, anti-ozonant, light stabilizer, the Lynn system peroxide decomposition agent, lubricant, paints, a foaming agent, a photo-setting resin, a thixotropic grant agent, etc. are raised.

An example of such an additive is indicated in each gazette of JP,4–69659,B, JP,7–108928,B, JP,63– 254149,A, and JP,64–22904,A, for example. Since a hardenability constituent of this invention has outstanding weatherability, heat resistance, water resisting property, and electric insulation resulting from a principal chain skeleton and it has high intensity and a physical-properties improvement offect of high elongation. It can use conveniently for electrical insulation materials, such as electric electronic component materials, such as a structural elastic sealing compound, a sealing material for multiple glass, a solar cell rear-face sealing agent, pre-insulation an electric wire, material for cables, a binder, adhesives, a paint, infusion, a coating material, a sealing agent for rust prevention / water proof, etc. It is useful especially when used for adhesives, a structural elastic sealing compound, or a sealing material for siding.

Example]

Although working example and a comparative example explain this invention concretely below, this invention is not limited to this.

(Working example 1-4, comparative example 1)

(A) As opposed to isobutylene system polymer (Kaneka Corp. make, trade name EP505S:isobutylene system polymer / paraffin series process oil = 100/50) 150 weight section which has a reactive silicon group as an ingredient, Epoxy resin (product [made from Oil recovery Shell Epoxy], trade name Epicoat 828) 5 weight section, Hydrogenation alpha-olefin oligomer (product [made from

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Idemitsu Petrochemistry], trade name PAO5004) 60 weight section, Colloid calcium carbonate (Maruo Calcium Co., Ltd make, trade name SHIRETTSU 200) 50 weight section, Golloid calcium carbonate (Maruo Calcium Co., Ltd make, trade name MC-5) 50 weight section, Heavy-calcium—carbonate (product [made from Shiroishi Calcium], trade name SOFUTON 3200) 40 weight section, Photo-setting resin (Toagosei make, trade name ARONIKKUSU M-309) 3 weight section, Benzotriazol system quiraviolet ray absorbent (Ciba-Geigy Japan make, trade name tinuin, 327) 1 weight section, Hindered phenolic artitoxidant (Sahkyo Co., Ltd. make, trade name SANORU LS-770) I weight section, hindered phenolic artitoxidant (Siba-Geigy Japan make, trade name IRUGA NOx 1010) 1 weight section, and water 5 weight section were measured respectively, and it often kneaded with a 3 paint roll, and was considered as base resin.

next — as the (B) ingredient — octylic acid (2-ethylhexanoic acid) calcium salt (the Nihon Kagaku Sangyo Co., Ltd. make.) The trade name NIKKAOKU Chicks Ca 5% toluene solution was used as a hardening agent [in / for the thing for which distearyl amine (the Kao Corp. make, Firmin D86) is respectively shown in Table 1 as a (C) ingredient and which could number/ of weight sections /— measure, could use the spatula, and was stirred and mixed / working example 1].

here — as the (B) ingredient of this invention — naphthenic acid vanadium (the Nihon Kagaku Sangyo Co., Ltd. make.) as a trade name NIKKANAFU textile V 2% toluene solution and the (C) ingredient — distearyl amine (the Kao Corp. make.) The number[of weight sections]—used thing which shows Firmin D86 in Table 1 respectively is made into working example 2, (B) as an ingredient — octylic acid (2-ethylhexanoic acid) iron (the Nihon Kagaku Sangyo Co., Ltd. make.) The number[of weight sections]—used thing which shows respectively lauryl amine (made by Wako Pure Chemical Industries, Ltd.) in Table 1 as a trade name NIKKAOKU Chicks Fe 6% toluene solution and a (C) ingredient is made into working example 3. The number[of weight sections]—used thing which 1 as a B ingredient as octylic acid (2-ethylhexanoic acid) titanium (3% toluene solution) and a (C) ingredient was made into working example 4. On the other hand, the number[of weight sections]—used thing which shows lauryl amine in Table 1 respectively as (B) octylic acid (2-ethylhexanoic acid) tin (Japanese east transformation Make, trade name neo SUTAN U-28) and a (C) ingredient was made into the comparative example 1.

Processor resin and hardening agents were mixed by the ratio shown in Table 1, respectively, and what constructed the aluminum base material to H type in accordance with the manufacturing method of the tensile adhesive property specimen to which this is specified JISA5758–1992 was filled up. Care-of-health conditions were made into 23 ***x7 +50 **x seven days. [per day] The used aluminum uses the alumite treatment aluminum of the 50x50x5-mm size based on JIS H4000 as a substrate, After purifying by methyl ethyl ketone (made by Wako Pure Chemical Industries, Ltd.) before being filled up with the compound, the primer (the Toray Industries Dow Coming make, trade mane D-2) was dried and used under ordinary temperature for spreading and 30 minutes.

In accordance with the tensile cement test method of 5758 – 1992 N of JIS A, the examination was presented with H type sample obtained by the above-mentioned method on condition of speed-of-testing 50 mm/min among 23 ** and the thermostatic chamber of 50**5% of humidity.

The recovery was measured by the following methods. First, after making H type sample which ended the above—mentioned care of health immersed into 50 ** warm water for one day and taking it out, it was neglected under the room temperature on the 1st. Subsequently, after having compressed and fixed to 70% so that sample thickness might be set to 12 to 8.4 mm, and heating a sample in 100 ** hot air drying equipment for one day, it released from compression and was neglected under the room temperature on the 1st. The thickness of the sample in this case was measured and the recovery to a compressed part was computed.

Time (leather-covered time) until it fills the same compound in an ointment can simultaneously as a measure which estimates hardenability and stretches a hide on the surface under 23 ** and the

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conditions of 50% of relative humidity was measured. The one where leather-covered time is shorter shows that hardenability is high. 0138]

A result is shown in Table 1. The filled hardened material is that cohesive failure is shown, and CF showing the fracture state in front is a fracture state which can be searched for as a sealing material.

Table 1]

	· · · · · · · · · · · · · · · · · · ·			実施例1	実施例2	実施例3	実施例4	比較例1
	7 4 3 -25 0	DDEAGG		天旭初山	3000 P12		<u> </u>	1047131
	(A) 成分	EP505S	phr			150		
		PAO5004	phr			60		
主		シーレッツ200	phr			50		
		ソフトン3200	phr	-		40		
		MC-5	phr			50		
		IL° J-1828	phr			5		
	İ	プロニックスM~309	phr			3		
		イルカ・ノックス1010	phr			1		
剤	İ	チヌピ ン327	phr			1		
	1	サノールLS-770	phr			1		
		水	phr			5		
	(B) 成分	オクチル酸Ca	phr	5.7				
硬		ナフテン酸V	phr		9.0			
`~	ł	オクチル酸Fe	phr			5.0		
化		オクチル酸Ti	phr				4.5	
'-	1	オクチル酸Sn	phr					3.0
剤	(C) 成分	ジステアリルアミン	phr	0.74	0.74			
777	(0, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	ラウリルアミン	phr	4		1.00	0.26	0.66
┢──	硬化性	皮張り時間	時間	4	4	>9	3	4
特	H型引張特性	M 5 0	MPa	0.05	0.07	0.08	0.08	0.1
l ' '		破断強度	MPa	0.45	0.64	0. 27	0.34	0.26
性		破断伸び	1 %	770	760	440	580	340
'-		破壊状態	<u></u>	CF	CF	ĊF	CF	CF
		復元率	1 %	57	73	27	29	54

When distearyl amine and lauryl amine were used together for octylic acid calcium, naphthenic acid Using an isobutylene system polymer as an ingredient (A) as shown in Table 1 as a (B) ingredient,

(working example 3) where used octylic acid iron as a (B) ingredient, and lauryl amine is used as a cowas fower than the comparative example 1, all had stability. High stability was shown when especially About stability, although the recovery at the time of using octylic acid titanium and octylic acid iron elongation after fracture to the comparative example 1 which used octylic acid tin and lauryl amine. catalyst, although it had been 9 hours or more leather-covered time, one day afterward, it checked covered time about equivalent has been secured as compared with the case where octylic acid tin and lauryl amine of the comparative example 1 are used together. On the other hand, in the case having hardened enough. The hardened material which was obtained also in the gap to use such carboxylic acid metal salt for (working example 1-4) showed breaking strength with high all, and vanadium, and octylic acid titanium as a co~catalyst (working example 1, 2, and 4), the leather naphthenic acid vanadium was used (working example 2).

and it kneaded once with 3 paint roll, and also checked that dried by decompression stirring at 120 ** weight section and a dripping inhibitor (made in Kusumoto Chemicals.) The amount part of trade name mixture might not involve in a bubble, and 1 liquid mold-curing nature constituent was obtained. What performed for 5 minutes. The cartridge made of paper (product made from the Showa round cylinder) CCR120 weight section and titanium oxide (Ishihara Sangyo Kaisha, Ltd. make, trade name TIPAQUE (C) The number of weight sections which shows lauryl amine (made by Wako Pure Chem) which is an cooling a mixture, it added and stirring mixing of the amount part of dehydrator (Nippon Unicar make, out for 15 minutes. The octylic acid zirconium (the Nihon Kagaku Sangyo Co., Ltd. make, trade name DISUPARON #6500 duplexs, benzotriazol system ultraviolet ray absorbent (Ciba-Geigy Japan make, trade name tinuvin 327) 1 weight section, and hindered amine light stabiliser (Sankyo Co., Ltd. make, trade name A-171) duplexs was carried out for 15 minutes. Next, it added and stirring mixing of the NIKKAOKU Chicks Zr.12% toluene solution) which is the (B) ingredient adjusted further beforehand, aminosilane coupling agent (Nippon Unicar make, trade name A~1120) 3 weight section was carried R-820) 20 weight section, and stirring mixing was carried out for 10 minutes. Took out the mixture, trade name SANORU LS-770) 1 weight section were measured, and it mixed in ******. the colloid stirring at 120 ** with 5L planetary mixer beforehand for 2 hours It mixed to trade name Hakuenka by which the coat was carried out was filled up with the aluminum for 1 liquid so that the obtained (A) As a polyether system polymer which has reactant silicon as an ingredient, As opposed to MS used the amount part of dibutyltin diacetyl acetonate (Japanese east transformation Make, trade calcium carbonate (the product made from Shiraishi Industry.) which dried this by decompression for 2 hours, and water content decreased to about 700 ppm with 5L planetary mixer again. After polymer S203 (made by Kaneka Corp.) 95 weight section, plasticizer (trade name PPG-3000) 55 names U-220) duplexs was made into the comparative example 2 instead of the mixture of said ingredient in Table 2, and the mixed thing were added, and decompression stirring mixing was (Working example 5, comparative example 2) octylic acid zirconium and lauryl amine.

aluminum of the 50x50x5-mm size based on JIS H4000, before the aluminum used as a substrate was examination was presented with H type sample obtained by the above-mentioned method on condition of for 23 **, and humidity 50 inside of **5% of thermostatic chamber and speed-of-testing health conditions were made into 23 **x14 +50 **x 14 days. [per day] Using the alumite treatment Industries, Ltd.). In accordance with the tensile cement test method of 5758 - 1992 N of JIS A, the method of the tensile adhesive property specimen specified to JIS A5758–1992 using the cancer of 00% 50%, respectively, and set TB and elongation at the time of a fracture to EB for the stress at marketing of 1 liquid mold-curing nature constituent from 1 liquid cartridge was filled up. Care-ofof 50 mm]/. The hardened material set stress when it elongated 150% to M50, M100, and M150 filled up with the compound, it was purified by methyl ethyl ketone (made by Wako Pure Chemical temperature for 24 hours or more, the specimen used for a tensile test was produced as follows. What constructed the aluminum base material to H type in accordance with the manufacturing After neglecting the above-mentioned 1 liquid mold-curing nature constituent under a room the time of a fracture.

Since an adhesive property turned into the important characteristic as a 1 liquid mold-curing nature constituent, the adhesive property was checked using anodized aluminium and a vinyl chloride steel 24/31 ページ

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with a 1 cm[, in width] x length of about 3 cm, and it was recuperated for 23 **x14 +50 ** x 14 days per day. The manual friction test was done after care of health, cutting an adhesion side deeply with a cutter knife, and the surface of the substrate was observed. The filled hardened material is that cohesive failure is shown, and OF in Table 2 is a fracture state searched for as a sealing material. nature constituent was placed from 1 liquid cartridge using commercial cancer to the crest shape plate. On each substrate which purified the surface by methyl ethyl ketone, 1 liquid mold-curing [0142]

Depths hardenability took out the portion which has hardened the surface which filled in the 80-mmlong polyethylene tube 1 liquid mold-curing nature constituent produced in working example 5, and was made at the tip of a tube in it at 16.5 mm in inside diameter on the day which the days to measure visited, and measured the hardened thickness with slide calipers.

cartridge is filled up with a constituent, it stored for two weeks in the hot air drying equipment which In order to check the practicality as a 1 liquid mold-curing nature constituent, where 1 liquid carried out the temperature control to 50 **, and the depths hardenability after storage was investigated. [0143]

[0144]

A result is shown in Table 2.

[0145]

Table 2

配合表	成分	製品名	実施例5	比較例2	
	(A)成分	\$203	95	95	施量重
	可塑剤	PPG3000	55	55	重量部
	炭酸カルシウム	白艶華CCR	120	120	商量重
	光安定剤	サノールLS-770	1	1	暗量重
	紫外線吸収剤	チヌピン327	1	1	第量部
	タレ防止剤	ディスパロン#6500	2	2	重量部
	酸化チタン	タイペークR-820	20	20	重量部
	脱水剤	A-171	2	2	重量部
	接着性付与剤	A-1120	3	3	哈量重
	(B)成分	オクチル酸ジルコニウム	8. 1		重量部
		U-220	Į	2	暗量重
	(C)成分	ラウリルアミン	1.4		重量部
H型引張機械特性		M50	0. 27	0. 19	MPa
	1	M100	0. 38	0. 31	MPa
		M150	C. 46	MPa	
	TE	3(破斷強度)	0. 84	0. 93	MPa
	E	3(破断伸び)	520	570	96
接着性	陽極直	変化アルミニウム	CF	CF	破壊状態
		塩ビ鋼鈑	CF	CF	破壊状態
硬化性	深部硬化性	23℃×1週間	7. 7	10.6	mm
		23℃×2週間	12. 2	15. 6	mm
貯蔵安定性	深部硬化性	23℃×1週間	7. 2	10.8	mm
(50℃×2週間貯蔵)		23℃×2週間	12. 0	16. 4	mm

working example 5, (B) By using the lauryl amine which acts the carboxylic acid zirconium salt which acts as a co-catalyst as a (C) ingredient further as a silanol condensation catalyst as an ingredient, As opposed to the polyether system polymer which serves as the (A) ingredient of this invention in adhesive property were revealed and it checked that 1 liquid mold-curing nature constituent which dibutyltin diacetyl acetonate as a tetravalent tin catalyst, almost comparable elongation and an Although hardenability was slightly inferior as compared with the comparative example 2 using may be sufficient for practical use was obtained (Table 2), (Synthetic example 1)

Use polyoxypropylene triol of the molecular weight 3,000 [about] as an initiator, and propylene oxide molecular weight about 26,000 (polystyrene reduced molecular weight in which the column measured the solvent using THF using the TOSOH TSK-GEL H type using TOSOH HLC-8120GPC as a liquidis polymerized in a zinc hexa oyanocobaltate glyme complex compound catalyst, Number average

sending system) polypropylene oxide was obtained. Then, the methanol solution of NaOMe of the equivalent was added 1.2 times to the hydroxyl group of this hydroxyl group end polypropylene oxide, and methanol was distilled off, and also the allyl chloride was added, and the hydroxyl group of the and was changed into the allyl group. By the above, the end obtained 3 organic—functions polypropylene oxide of the number average molecular weight 26,000 [about] which is an allyl group. [0] 46]

introduction rate was measured by the following methods by ¹H-NMR (it measures in a CDC)₃ solvent azeotropically at 90 ** and distilling off hexane under decompression. On the other hand, after adding reaction. The relative value of the peak integral value of the proton (near CH₃(CH₃O) ₂Si-CH₂-CH₂-0.6ppm) of the methylene group combined with the silicon atom of the end silyl group: Two ** et al., conversion) I, 7.0 g of dimethoxymethylsilane was dropped. After making the mixed solution react at 30 ** for 2 hours, unreacted dimethoxymethylsilane was distilled off under decompression and the 30micro of platinum divinyl disiloxane complex (it is 3% of the weight of xylene solution by platinum polypropylene oxide main chain of said allyl end 3 organio-functions polypropylene oxide before a using JEOL JNM-LA400). Relative-value:1 of the peak integral value of the allyl end proton (near polypropylene oxide and 10 g of hexane which were obtained above to 1L autoclave, dehydrating hydrosilylation reaction . Receive the peak integral value of the ${\sf CH}_3$ group (near 1.2 ppm) of the polypropylene oxide main chain of the silyl end polypropylene oxide (A-1) after a hydrosilylation reactive silicon group content polyoxyalkylene series polymer (A-1) was obtained. The number The nitrogen purge was carried out, after having added 500 g of allyl end 3 organic-functions $\mathrm{CH_2}$ =CH-CH₂=:5.1 ppm) to the peak integral value of the CH₃ group (near 1.2 ppm) of the average molecular weight of the obtained polymer (A-1) was about 26,000. The silyl group a silyl group introduction rate (**2.) /**1. It was 78%

After attaching a three-way cook to the resisting pressure glass container of 2L and carrying out the nitrogen purge of the inside of a container to it. Using an injector, in a container 262.5 ml of ethylosclohexane (thing dry by neglecting 1 or more night with the molecular sieves 3A), and 787.5 ml of toluene (thing dry by neglecting 1 or more night with the molecular sieves 3A), p-DCC (the collowing compound (A)) 4.85g (21.0mmol) was added.

0147] Formula 11]

Next, the resisting pressure glass liquefied gas extraction pipe with a needle valve containing 438 ml (5.15 mol) of isobutylene monomers was connected to the three—way cock, and after attaching the polymerization vessel all over dry ice / ethanol bus of -70 ** and cooling, the inside of a container was made decompression using the vacuum pump. After opening the needle valve and introducing an isobutylene monomer in a polymerization vessel from a liquefied gas extraction pipe, the inside of a container was returned to ordinary pressure by introducing nitrogen from one side in a three-way cock. Next, 0.72 g (7.7mmol) of 2-methylpyridine was added. Next, 10.58 ml (96.5mmol) of titanium tetrachloride was added, and the polymerization was started. No minutes after the polymerization start, 7.20 g (63.0mmol) of allyl timethylsilane was added, and the introduction reaction of the allyl egroup was performed to molecular chain terminals. After adding allyl trimethylsilane and 200 ml of water washed [120 minutes] the reaction solution 4 times, the allyl end isobutylene system polymer was obtained by distilling off a solvent.

to 40.5 Subsequently, the allyl end isobutylene system polymer 200g obtained in this way, the paraffin-base process oil (the Idemitsu Kosan make.) which is a hydrocarbon system plasticizer After mixing trade nttp://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http%3A%2F%2Fwww4.ipdl.i.. 2010/05/06

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name Diana process PS-32 100g and carrying out temperature up to about 75 **, the 7.5×10⁻⁵ equivalent was added [methyl dimethoxysilane] for 2.4 Eq and a platinum (vinyl siloxane) complex to the allyl group of an end, and the hydrosilylation reaction was performed. FT-IR (Shimadzu IR-408) performed reaction pursuit, and the absorption based on the olefin of 1640 cm⁻¹ disappeared in about 20 hours.

[0149]

2/1 of PS-32 which is the isobutylene system polymer and plasticizer which have a reactive silicon group in the both ends of the target chain of the mixtures (A-2) of the weight ratio were obtained. In sol

In this way, if the obtained polymer is measured by the GPC method (it was considered as the liquid-sending system and, as for the solvent, the column used CHCl₃ using Shodex K-804 using LGModule1 made from Waters). The number average molecular weight was 17,600 and Mw/Mn (ratio of weight average molecular weight to a number average molecular weight) was 1.23. ¹H-NMR (using VarianGemini300) The proton which belongs to each structure by measurement in CDCl₃ (the proton of initiator origin: 6.5–7.5 ppm) The methyl proton combined with the silicon atom of polymer-terminal origin: As a result of measuring and measuring the intensity of 0.0–0.1 ppm and the resonance signal of methoxy proton:3.4 – 3.5, the end silyf functional group number Fn (number of the silyl group per one molecule of isobutylene polymer) was 1.76.

(Working example 6–15, comparative example 3)

(A) As an ingredient, various additive agents were measured, respectively, and it often kneaded with a 3 paint roll according to the formula shown in Table 2, using the reactive silicon group content polyoxyalkylene series polymer (A–1) obtained in the synthetic example 1, and was considered as base resin.

[0151]

Next, as a (B) ingredient which is a silanol condensation catalyst, lauryl amine was further measured for the various carboxylic acid metal salt shown in Table 1 to the above—mentioned base resin as a (C) ingredient, the spatula was used for it, and it stirred and mixed for 3 minutes. It was made for all the numbers of mols of the metal atom containing the number of addition parts of the various carboxylic acid metal salt of the (B) ingredient to become the same here. The mold about 3 mm thick was used and filled up with the spatula after mixing, and care of health on 23 **x3 +50 **x the 4th sapecimed. I per day I From the obtained hardened material sheet, the No. 3 type dumbbell specimen specified to JIS K 6301 was pierced, and the tensile test was carried out in the autograph (tension speed 200 mm/min). The modulus (M50) in the time of being extended 50%, the intensity at the time of a dumbbell fracture (Eb) were measured.

10 132.]
The combination presentation of base resin, a curing catalyst, etc. and the physical-properties evaluation result of the hardened material obtained from it are shown in Table 3.

Fig. 13.2.

Table (

	祖成(重量部)	実施例										(比較)			
	6	7	. 8	9	10	11	12	13	14		3				
(A) 成分	成分 A-1		95	95	95	95	95	95		95	95	95	95		
	白野華CCR ⁽³⁾	白石工業(株)	.60	. 60	60	60	69	60	60	60	60	60	60		
充填材	£" 335/18 ⁽⁰⁾	白石工業(株)	60	60	69	60	60	60	60	60	60	60	60		
	\$7(}\SB ⁽¹⁾	白石がかん(株)	20	20	20	20	50	20	20	20	20	20	20		
可塑劑	7クトコール P-23(3) 武田南品工業 (株)		60	60	60	60	60	60	60	60	60	60	60		
クソ性付与剤	ディスハ ロン ま305 ⁽⁴⁾	補本化成(株)	3	1	3	3	3	3	3	3	3	3	3		
表面改貨劑	701-57XH-309(F)	東亜会成化学(株)	3	1	3	3	3	3	3	3	3	3	3		
外棋吸収剂	53t >327 ⁽⁰⁾	テバガイギー(株)	1	_	1			_1_	_	1	. 1	1	1		
酸化防止剂	483 79921010 ⁽¹⁾	チバガイギー (株)	1	1	_	1		1		ı	1	1	1_1		
(B) 成分	こったようチックス家共 GX (T) ^(a)	日本化学産業(株)	6, 6												
	ナフテックス鉄5%(T) ^(E)	日本化学産業(株)		7.9							i		L		
	2 - エチルヘキサン	食チタニウム ⁽¹⁰⁾			11										
	ナンチックスパ ナシ ウム2%(P) ⁽¹¹⁾	日本化学産業(株)				3.8						L	<u> </u>		
	こっちょうチェクスカルシウム5%(T) ⁽¹³⁾	日本化学産業(株)					5.6						I		
	プキャット15 G ^(M)	日本化学産業(株)						1,0					<u> </u>		
	こっカオクチックスパ リウム8%(P) (74)	日本化学産業(件)							12						
	こっちオクチックスマンカーン8%(T)(140	. 日本化学 双 篇(株).								4.9					
	こっきオクチックスニックル6% (P) ⁽¹⁴⁾	日本化学産業(株)									5.8		L		
	ニッカキウチックシ ホコニウ&12%(T) ⁽¹¹⁾	日本化学産業(株)										5.4			
	2 - エチルヘキサ	ン数スズ(15)													
(C) 成分	ラウリルフ	'ミン	0.66	0.56	0.66	0.66	0.65	0.66	0.66		0.66	0.56	0.0		
更化物物性	M50	(NPa)	0. 19	0.18	0.21	0.18	0.13	0.12	0.12	0.16	0.10 1.20	0.20	<u>.</u>		
	Tb Eb	(MPa) (%)	0. 63 650	0.75 640	7.50	1,12 520	0.94 860	760	0.92 810	0. 99 770	1100	1.27 890	. O. 1		

When the various carboxylic acid metal salt of working example 6–15 is used compared with the comparative example 3 using 2-ethylhexanoic acid tin as a silanol condensation catalyst as shown in Table 3, the value of Tb (breaking strength) and Eb (elongation after fracture) of hardened material physical properties is larger.

High elongation and high intensity were shown.

(Working example 16-28, comparative example 4)

(A) Using the mixture (A-2) of the isobutylene system polymer and plasticizer which have as an ingredient the reactive silicon group obtained in the synthetic example 2, various additive agents were measured, respectively, and it often kneaded with a 3 paint roll according to the formula shown in Table 4, and was considered as base resin.

Next, the various carboxylic acid metal salt shown in Table 4 as a (B) ingredient which is a silanol

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condensation catalyst was measured, and also as a (C) ingredient, lauryl amine was used together to the above-mentioned base resin, the spatula was used by working example 16-28 and the comparative example 4 for it, and it stirred and mixed for 3 minutes to it. It was recuperated at 23 ** after mixing, and the surface hardened state was evaluated five days afterward. It was made for all the numbers of mols of the metal atom containing the number of addition parts of the various carboxylic acid metal salt of the (B) ingredient to become the same here.

[0155]
The evaluation result of the hardened state five days after the combination presentation of base resin, a curing catalyst, etc. is shown in Table 4. What the surface had hardened O in front for five days afterward is shown, and it is shown that x had not hardened five days after.

[0156] [Table 4]

									-m&				_			比较例
組成 (重要部)			ļ	17"	18	19	żū	21	実施例 22 1	23	24	25	26	27	28	4
	A - 2		150	158	150	180	150	150	150	150	150	150	150	150	[50	150
(A) 成分	÷-1-77200 ⁽¹⁾	丸属ががな (株)	50	50	50	50	50	50	50	50	50	50	50	50	50	50
	9-977230 RC-5 ⁽¹⁾	九馬胡沙丘(株)	50	50	50	50	50	50	50	50	50	50	50	50	50	50
充填材	771-73508 GB	白石がかりに(株)	40	40	40	40	40	40	40	40	40	40	40	40	40	40
	PA05004 ⁽¹⁾	出光興産(株)	60	60	£0	60	50	60	60	60	60	60	69	60	60	60 .
<u>可型剤</u> エポキシ樹脂	1t' 3-1828(0)	治化シェルは、キン(株)	5	5	5	5	5	5	5	5	5	5	- 5	5	- 5	5
	702+724-308 ^(b)	東亞合成化学(株)	1 1	3	3	3	3	- 3	3	3	3	1	3	3	3	3_
表面改質剤	53t' 7427' ³⁰	チバガイギー(株)	1	· ·		ī	Ť	1			i		1	1	1	١,
<u> </u>	432 /2221010 ⁽⁷⁾	チバガイギー(株)		-	1	i				1	1	1	1		- 1	1
酸化防止剂	#J-MLS-770 ⁽⁴⁾	三世(株)	┼-			+		1	- 1	1	i	1	1		1	1
光安定割	37-MC3-110.	<u>二大 (律)</u>		1 5	 	5	5	- 5	5	- 5	5	5	5	5	5	S
-	a sale brancacas (D)	日本化学産業(排)	6.6	 									6.6			
(D) AXX	ニッカオクチックス 飲らな(工) ***		1 2.0	7.9												
	ナフテックス鉄55 (T) (III 日本化学産業(株)		1-	1.53	11							-		11		
	2 - エチルヘキサン闘デタニウム ⁽⁽⁾⁾ ナフテックスパナジ ウム28 (P) ⁽¹⁾ 日本化学座業(株)			<u> </u>		18	_	_					-		7.8	1
	こったオクチックスカルシウム5%(T) (F)	日本化学産業(株)			_	<u> </u>	5.6									
	27877779X8477551()	日本化学機能(株)	-	 		_		12					Ì			
	278727733A 1/3483 (P)	日本化学産業(株)	! 		-				4.9				1			$\overline{}$
	ニッカオクチックスマンカ ン名葉(7) (15) ニッカオクチックスニックルラキ(P) (14)	日本化学産業(株)								6.8					-	
		日本化学産業(株)	-		-					***	3.4					
			—		-						W. 7	5.4				
	=9849f-997 &1=26128 (T) (10)	日本化学産業(株)	! 		_		-						 		T	2.5
*****	こったオクチックス型船181x ^{((の)} ラウリルフ	日本化学座館(株)	0.66	0.66	33.0	0.56	0.68	0.55	0.66	0.66	0.66	0.66	 	-		6.66
"(C) 成分	5日後の神化状態	37	0.00	0	()	0	0		0	Ö	0	O	10	I. O	O	X.
(5): トリメテロ (5): 2 - (3, 1) (5): 2 - (3, 2) (8): ピス (2, 3): 2 - エデル (10): ナフ・エテンチ (11): 2 - フ・エチチ (14): 2 - エエチチ (15): 2 - エエチチ (16): 2 - エエチチ (18): 2 - エエチチ	ルシウム	ドロキシーン・(イー・(イー・) (イー・(イー・) (イー・(イー・) (イー・) (イー	チルーキ ランス	ー : 6 % 3 ** ** ** ** ** ** ** ** ** ** ** ** *	キシフ: E:5% 8% 12%	:=./v) :	プロピオ	≵− ⊦]								

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spite of having used the silanol condensation catalyst of the non-tin series, the surface has hardened When the various carboxylic acid metal salt of working example 16-28 is used as shown in Table 4, in

Practical hardenability was shown.

On the other hand, when the carboxylic acid zinc of the comparative example 4 was used,

nardenability had not hardened five days after bad.

[0157]

carboxylic acid zirconium, and a hardenability [** and others] constituent, In spite of being a non-tin molecule, (B) Carboxylic acid calcium, carboxylic acid vanadium, carboxylic acid iron, Carboxylic acid titanium, carboxylic acid potassium, carboxylic acid barium, Carboxylic acid manganese, carboxylic As mentioned above, the organic polymer which has at least one reactive silicon group in the (A) acid nickel, carboxylic acid cobalt, one or more sorts of carboxylic acid metal salt chosen from a series catalyst, practical hardenability is shown, and it tums out that hardened material physical properties are also fitness (high elongation and high intensity)

(Constituent using the trimethoxysilyl group as a reactive silicon group)

組成(重量部)

ニッカオクチックスカルシウム5%(T)

terminals as a reactive silicon group, a cure rate will become large from the polymer of the synthetic For example, the same hardenability constituent as working example 6-15 can be prepared using the polymer indicated for the example 1 of manufacture of JP,H11~12480,A. The tack free time of this methyl dimethoxy silyl group. Such a polymer is indicated to JP,H11-12480,A or JP,2001-72855,A. example 6–15 can be prepared using the polymer indicated for the example 1–4 of manufacture of constituent is shorter than working example 6-15. The same hardenability constituent as working If a trimethoxysilyl group uses the polyoxyalkylene series polymer which exists in molecular chain Therefore, a catalyst amount can be lessened, if cure time is made into the same grade and the JP,2001-72855,A. The tack free time of this constituent is shorter than working example 6-15. example 1. The direction of a trimethoxysilyl group of this is because reactivity is larger than a polymer which has the above-mentioned trimethoxysilyl group will be used. 0158

if the mixture of the polymer which has a trimethoxysilyl group, and the polymer which has a methyl material are freely controllable. For example, the hardenability constituent of working example 6-15 oan be prepared using the polymer which mixed the polymer which has a trimethoxysilyl group, and dimethoxy silyl group is used as a polymer, cure time and the physical properties of a hardened the polymer which has a methyl dimethoxy silyl group by the weight ratio of 1:10-10:1.

(Synthetic example 3)

Working example when the polymer which has a trimethoxysilyl group is used is shown below

実施例

100

6.6

29

100

6.6

number average molecular weight 17,000 and the polyoxypropylene diol 1000g of molecular-weightdistribution Mw/Mn=1.20 as 19g of gamma-isocyanate propyltrimethoxysilane (Nippon Unicar make compound catalyst. It polymerizes. The obtained polypropylene oxide. Use and to the compound The inside of bottom of N, atmosphere 1L autoclave, According to a composite metal complex Y-5187), and a catalyst, 0.05 g of dibutyl tin screw isooctylthioglycolate (made in [U-360]

polyoxypropylene (A-3) by which the trimethoxysilyl group was introduced into about 80% of ends was longer detected in IR under the nitrogen air current in this, and the reactive silicon group content transformation [Japanese east]) is added, It reacted at 90 ** until the isocyanate group was no

obtained. (Working example 29–32)

ngredients according to the formula shown in Table 5 using the polyoxyalkylene series polymer (A-3) ngredient was measured, respectively, the spatula was used, and it stirred and mixed for 30 seconds. it was made for all the numbers of mols of the metal atom containing the number of addition parts of which has the trimethoxysilyl group obtained in the synthetic example 3, (C) Lauryl amine which is an surface was lightly pressed down with the spatula after mixing, and time (leather-covered time) until (A) The polyoxyalkylene series polymer (A-1) which has as an ingredient the methyl dimethoxy silyl group obtained in the synthetic example 1, The various carboxylic acid metal salt which is the (B) the various carboxylic acid metal salt of the (B) ingredient to become almost the same here. The constituent stops adhering at the tip of a spatula was measured. http://www4.ipdl.inpit.go.jp/ogi-bin/tran_web_ogi_ejje?atw_u=http%3A%2F%2Fwww4.ipdf.i.. 2010/05/06

An evaluation result is shown in Table [0161]

Table 5

成分 (A)

成分

成分 ラウリルアミン 0.66 (C) 0.66 300min以上 (23°C) 皮張り時間 41min 23min エチルヘキサン酸鉄のミネラルスピリット溶液、金属(Fe) 含有量 キサン酸カルシウムのミネラルスピリット溶液、金属(Ca) 含有量:5%

日本化学産業

A - 1

As shown in Table 5, the polyoxyalkylene series polymer (A-3: working example 30 and 32) which has a trimethoxysilyl group had the cure rate quicker than the polyoxyalkylene series polymer (A-1:

32

100

5.6

0.66

1**05m**in

31

100

5.6

0.66

working example 29 and 31) which has a methyl dimethoxy silyi group.

polyoxyalkylene series polymer (A-3) which has the trimethoxysilyl group obtained in the synthetic example 3. The tack free time of this constituent has short cure time, and good physical properties The same hardenability constituent as working example 6–15 can be prepared using the

(Constituent which used the epoxy resin together)

The same hardenability constituent as working example 6-15 can be prepared using the constituent which used together reactive silicon group containing organic polymer and an epoxy resin. When this constituent is used, the manifestation of adhesive strength is quick.

[0163] [Effect of the Invention]

mechanical physical property which has high intensity and high elongation as compared with the case where the carboxylic acid tin currently used conventionally is used as a curing catalyst. Therefore, the hardenability constituent of this invention is very useful as various elastomers, such as a sealing titanium, carboxylic acid potassium, carboxylic acid barium, The hardenability constituent containing revealing practical hardenability and stability, the hardened material obtained shows the outstanding atom, and has at least one silicon content group which can construct a bridge by forming a siloxane (A) The organic polymer which has the hydroxyl group or hydrolytic basis combined with the silicon bond, (B) Carboxylic acid calcium, carboxylic acid vanadium, carboxylic acid iron, Carboxylic acid one or more sorts of carboxylic acid metal salt which it comes to choose out of carboxylic acid manganese, carboxylic acid nickel, carboxylic acid cobalt, and a carboxylic acid zirconium, While material, adhesives, a binder.

[Translation done.]